```
FILE 'REGISTRY' ENTERED AT 10:33:28 ON 01 OCT 2009
               STRUCTURE UPLOADED
L2
              0 S L1
L3
              STRUCTURE UPLOADED
L4
             0 S L3
L5
              STRUCTURE UPLOADED
L6
             0 S L5
L7
             2 S L5 SSS FULL
L8
               STRUCTURE UPLOADED
L9
             0 S L8
L10
             0 S L8 SSS FULL
     FILE 'STNGUIDE' ENTERED AT 10:38:12 ON 01 OCT 2009
     FILE 'HCAPLUS' ENTERED AT 10:43:33 ON 01 OCT 2009
             0 S (LYSOPHOSPHTIDIC ACID) (4A) (INHIB? OR ANTAG?)
L12
         717126 S PHOSPHATE OR PHROPHOSPHATE OR PHOSPHORIC
L13
        220274 S CHOLESTEROL OR HYPERCHOLESTEROL? OR HYPERLIPID? OR ATEHROSCLE
L14
             0 S L11 AND L12
L15
             0 S L11 AND L13
L16
          6976 S L12 AND L13
     FILE 'STNGUIDE' ENTERED AT 10:43:40 ON 01 OCT 2009
     FILE 'HCAPLUS' ENTERED AT 10:44:18 ON 01 OCT 2009
L17
            314 S (LYSOPHOSPHATIDIC ACID) (4A) (INHIB? OR ANTAG?)
         744163 S PHOSPHATE OR PYROPHOSPHATE OR PHOSPHORIC
T.18
        220274 S CHOLESTEROL OR HYPERCHOLESTEROL? OR HYPERLIPID? OR ATEHROSCLE
L19
L20
            62 S L17 AND L18
             3 S L17 AND L19
L21
L22
          7389 S L18 AND L19
     FILE 'REGISTRY' ENTERED AT 15:38:02 ON 01 OCT 2009
                EXP SERINE PHOS/CN
                EXP SERINE PHOSPHATE/CN
L1
              1 S E5
                EXP SERINE PHOSPHORIC/CN
     FILE 'HCAPLUS' ENTERED AT 15:38:45 ON 01 OCT 2009
L2
             67 S L1/THU
L3
         194608 S NEOINTIM? OR ATHEROSCLEROSIS OR STENT OR CARDIOVASCULAR
L4
             0 S L2 AND L3
L5
             31 S L2 AND (PY<2003 OR AY<2003 OR PRY<2003)
```

=> file registry COST IN U.S. DOLLARS

FULL ESTIMATED COST

SINCE FILE TOTAL ENTRY SESSION

0.22

0.22 FILE 'REGISTRY' ENTERED AT 10:33:28 ON 01 OCT 2009

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Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 29 SEP 2009 HIGHEST RN 1186580-18-6 DICTIONARY FILE UPDATES: 29 SEP 2009 HIGHEST RN 1186580-18-6

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH June 26, 2009.

Please note that search-term pricing does apply when conducting SmartSELECT searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

Uploading C:\Program Files\STNEXP\Queries\10821739monophosphate.str

```
chain nodes :
1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
24 25 30 31 32 33 34 35 36 37 38 39 41 42 43 44 45 46 47 48 49
50 51 52 53
54 55 58
          59
              60 61
                     62 63 64 66 67 68 69 70 71
                                                      72
                                                          73
                                                                 75 76 77
78 79 80
81 82 83 84 85 86
chain bonds :
1-2 1-3 1-4 1-5 5-30 6-8 6-14 7-10 7-15 8-9 8-12 9-13 10-11 10-16
11-17
13-18 14-19 15-20 16-21 19-22 19-24 21-23 21-25 31-32 31-34 32-33 32-35
33-36 34-37
35-41 37-38 37-39 42-44 42-50 43-46 43-51 44-45 44-48 45-49 46-47 46-52
47-53 50-54
52-55 58-59 58-61 59-60 59-62 60-63 61-64 66-67 66-68 66-69 67-71 67-72
68-70 73-74
73-75 73-76 74-78 74-79 75-77 80-81 80-82 80-83 81-85 81-86 82-84
exact/norm bonds :
1-2 1-3 1-4 1-5 5-30 8-12 10-16 13-18 14-19 15-20 16-21 19-22 19-24
21-23 21-25 32-35 34-37 35-41 37-38 37-39 44-48 46-52 50-54 52-55 59-62
61-64 66-69 67-71
67-72 73-76 74-78 74-79 80-83 81-85 81-86
exact bonds :
6-8 \quad 6-14 \quad 7-10 \quad 7-15 \quad 8-9 \quad 9-13 \quad 10-11 \quad 11-17 \quad 31-32 \quad 31-34 \quad 32-33 \quad 33-36 \quad 42-44
```

42-50 43-46 43-51 44-45 45-49 46-47 47-53 58-59 58-61 59-60 60-63 66-67

66-68 68-70

G1:[*1],[*2]

```
Connectivity:
```

24:1 X maximum RC ring/chain 25:1 X maximum RC ring/chain

Match level : 1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS

10:CLASS 11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS 18:CLASS 19:CLASS 20:CLASS 21:CLASS 22:CLASS 23:CLASS 24:CLASS 25:CLASS 30:CLASS 31:CLASS

32:CLASS 33:CLASS

34:CLASS 35:CLASS 36:CLASS 37:CLASS 38:CLASS 39:CLASS 41:CLASS 42:CLASS 43:CLASS 44:CLASS

45:CLASS 46:CLASS 47:CLASS 48:CLASS 49:CLASS 50:CLASS 51:CLASS 52:CLASS 53:CLASS 54:CLASS

55:CLASS 58:CLASS 59:CLASS 60:CLASS 61:CLASS 62:CLASS 63:CLASS 64:CLASS 66:CLASS 67:CLASS

68:CLASS 69:CLASS 70:CLASS 71:CLASS 72:CLASS 73:CLASS 74:CLASS 75:CLASS 76:CLASS 77:CLASS 78:CLASS 79:CLASS 80:CLASS 81:CLASS 82:CLASS 83:CLASS 84:CLASS 85:CLASS

86:CLASS

L1 STRUCTURE UPLOADED

=> d 11 L1 HAS NO ANSWERS

L1 STR

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

Structure attributes must be viewed using STN Express query preparation.

=> 8 11

SAMPLE SEARCH INITIATED 10:33:56 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED -

0 ITERATIONS

0 TO ITERATE

100.0% PROCESSED SEARCH TIME: 00.00.01

0 ANSWERS

FULL FILE PROJECTIONS: ONLINE **COMPLETE** BATCH **COMPLETE**

PROJECTED ITERATIONS: PROJECTED ANSWERS:

0 TO 0 0 TO Λ

L2

0 SEA SSS SAM L1

Uploading C:\Program Files\STNEXP\Oueries\10821739monophosphate2.str

```
1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
24 25 30 31 32 33 34 35 36 37 38 39 41 42 43 44 45
                                                         46
50 51 52 53
54 55 58
         59 60 61 62 63 64 66 67 68 69 70 71 72 73 74 75 76 77
78 79 80
81 82 83 84 85 86
chain bonds :
1-2 1-3 1-4 1-5 5-30 6-8 6-14 7-10 7-15 8-9 8-12 9-13 10-11 10-16
11-17
13-18 14-19 15-20 16-21 19-22 19-24 21-23 21-25 31-32 31-34 32-33 32-35
33-36 34-37
35-41 37-38 37-39 42-44 42-50 43-46 43-51 44-45 44-48 45-49 46-47 46-52
47-53
     50-54
52-55 58-59 58-61 59-60 59-62 60-63 61-64 66-67 66-68 66-69 67-71 67-72
68-70 73-74
73-75 73-76 74-78 74-79 75-77 80-81 80-82 80-83 81-85 81-86 82-84
exact/norm bonds :
1-2 1-3 1-4 1-5 5-30 8-12 10-16 13-18 14-19 15-20 16-21 19-22 19-24
21-23 21-25 32-35 34-37 35-41 37-38 37-39 44-48 46-52 50-54 52-55 59-62
61-64 66-69 67-71
67-72 73-76 74-78 74-79 80-83 81-85 81-86
```

6-8 6-14 7-10 7-15 8-9 9-13 10-11 11-17 31-32 31-34 32-33 33-36 42-44

chain nodes :

exact bonds :

```
42-50 43-46 43-51 44-45 45-49 46-47 47-53 58-59 58-61 59-60 60-63 66-67
66-68 68-70
73-74 73-75 75-77 80-81 80-82 82-84
G1:[*1],[*2],[*3],[*4],[*5],[*6],[*7],[*8],[*9]
Connectivity:
24:1 X maximum RC ring/chain 25:1 X maximum RC ring/chain
Match level :
1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS
10:CLASS 11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS
18:CLASS 19:CLASS
20:CLASS 21:CLASS 22:CLASS 23:CLASS 24:CLASS 25:CLASS 30:CLASS 31:CLASS
32:CLASS 33:CLASS
34:CLASS 35:CLASS 36:CLASS 37:CLASS 38:CLASS 39:CLASS 41:CLASS 42:CLASS
43:CLASS 44:CLASS
45:CLASS 46:CLASS 47:CLASS 48:CLASS 49:CLASS 50:CLASS 51:CLASS 52:CLASS
53:CLASS 54:CLASS
55:CLASS 58:CLASS 59:CLASS 60:CLASS 61:CLASS 62:CLASS 63:CLASS 64:CLASS
66:CLASS 67:CLASS
68:CLASS 69:CLASS 70:CLASS 71:CLASS 72:CLASS 73:CLASS 74:CLASS 75:CLASS
76:CLASS 77:CLASS
78:CLASS 79:CLASS 80:CLASS 81:CLASS 82:CLASS 83:CLASS 84:CLASS 85:CLASS
86:CLASS
L3
      STRUCTURE UPLOADED
=> s 13
SAMPLE SEARCH INITIATED 10:35:02 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED -
                                    1 TO ITERATE
```

0 ANSWERS

1 ITERATIONS

COMPLETE

8.0

0

1 TO

0 TO

Uploading C:\Program Files\STNEXP\Queries\10821739monophosphate3.str

FULL FILE PROJECTIONS: ONLINE **COMPLETE** BATCH

0 SEA SSS SAM L3

100.0% PROCESSED

SEARCH TIME: 00.00.01

PROJECTED ITERATIONS:

PROJECTED ANSWERS:

L4

```
1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
24 25 30 31 32 33 34 35 36 37 38 39 41 42 43 44 45
                                                         46
50 51 52 53
54 55 58
         59 60 61 62 63 64 66 67 68 69 70 71 72 73 74 75 76 77
78 79 80
81 82 83 84 85 86
chain bonds :
1-2 1-3 1-4 1-5 5-30 6-8 6-14 7-10 7-15 8-9 8-12 9-13 10-11 10-16
11-17
13-18 14-19 15-20 16-21 19-22 19-24 21-23 21-25 31-32 31-34 32-33 32-35
33-36 34-37
35-41 37-38 37-39 42-44 42-50 43-46 43-51 44-45 44-48 45-49 46-47 46-52
47-53
     50-54
52-55 58-59 58-61 59-60 59-62 60-63 61-64 66-67 66-68 66-69 67-71 67-72
68-70 73-74
73-75 73-76 74-78 74-79 75-77 80-81 80-82 80-83 81-85 81-86 82-84
exact/norm bonds :
1-2 1-3 1-4 1-5 5-30 8-12 10-16 13-18 14-19 15-20 16-21 19-22 19-24
21-23 21-25 32-35 34-37 35-41 37-38 37-39 44-48 46-52 50-54 52-55 59-62
61-64 66-69 67-71
67-72 73-76 74-78 74-79 80-83 81-85 81-86
```

6-8 6-14 7-10 7-15 8-9 9-13 10-11 11-17 31-32 31-34 32-33 33-36 42-44

chain nodes :

exact bonds :

```
42 - 50 \quad 43 - 46 \quad 43 - 51 \quad 44 - 45 \quad 45 - 49 \quad 46 - 47 \quad 47 - 53 \quad 58 - 59 \quad 58 - 61 \quad 59 - 60 \quad 60 - 63 \quad 66 - 67 \quad 60 - 63 \quad 60 -
66-68 68-70
73-74 73-75 75-77 80-81 80-82 82-84
G1:[*1],[*2],[*3],[*4],[*5],[*6],[*7],[*8],[*9]
Connectivity:
24:1 X maximum RC ring/chain 25:1 X maximum RC ring/chain 54:1 X maximum RC
ring/chain
55:1 X maximum RC ring/chain 64:1 X maximum RC ring/chain
Match level :
1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS
10:CLASS 11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS
18:CLASS 19:CLASS
20:CLASS 21:CLASS 22:CLASS 23:CLASS 24:CLASS 25:CLASS 30:CLASS 31:CLASS
32:CLASS 33:CLASS
34:CLASS 35:CLASS 36:CLASS 37:CLASS 38:CLASS 39:CLASS 41:CLASS 42:CLASS
43:CLASS 44:CLASS
 45:CLASS 46:CLASS 47:CLASS 48:CLASS 49:CLASS 50:CLASS 51:CLASS 52:CLASS
53:CLASS 54:CLASS
 55:CLASS 58:CLASS 59:CLASS 60:CLASS 61:CLASS 62:CLASS 63:CLASS 64:CLASS
66:CLASS 67:CLASS
68:CLASS 69:CLASS 70:CLASS 71:CLASS 72:CLASS 73:CLASS 74:CLASS 75:CLASS
76:CLASS 77:CLASS
 78:CLASS 79:CLASS 80:CLASS 81:CLASS 82:CLASS 83:CLASS 84:CLASS 85:CLASS
86:CLASS
L5 STRUCTURE UPLOADED
=> s 15
SAMPLE SEARCH INITIATED 10:36:24 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED -
                                                                                                      1 TO ITERATE
100.0% PROCESSED
                                                              1 ITERATIONS
                                                                                                                                                                          0 ANSWERS
SEARCH TIME: 00.00.01
FULL FILE PROJECTIONS: ONLINE **COMPLETE**
                                                               BATCH **COMPLETE**
PROJECTED ITERATIONS:
                                                                                     1 TO
                                                                                                             80
PROJECTED ANSWERS:
                                                                                      O TO
                                                                                                                   0
1.6
                                    O SEA SSS SAM L5
=> s 15 sss full
FULL SEARCH INITIATED 10:36:30 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED -
                                                                                               23 TO ITERATE
100.0% PROCESSED
                                                            23 ITERATIONS
                                                                                                                                                                          2 ANSWERS
SEARCH TIME: 00.00.01
                                  2 SEA SSS FUL L5
L7
=> d 17 scan
```

L7 2 ANSWERS REGISTRY COPYRIGHT 2009 ACS on STN IN Hexadecanoic acid, (1S)-1-[[[[([S)-1-carboxy-2-hydroxyethyl]amino]oxy]hydroxyphosphinyl]oxy]methyl]-1,2-ethanediyl ester

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

- 2 ANSWERS REGISTRY COPYRIGHT 2009 ACS on STN L-Alanine, 3-(phosphonodioxy)-
- IN
- C3 H8 N O7 P

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

ALL ANSWERS HAVE BEEN SCANNED

Uploading C:\Program Files\STNEXP\Queries\10821739pyrophosphate.str

G1:[*1],[*2]

chain nodes :

Connectivity :

- 30:1 X maximum RC ring/chain 31:1 X maximum RC ring/chain 32:1 X maximum RC ring/chain
- 33:1 X maximum RC ring/chain
- Match level:
 1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS
 10:CLASS 11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS
 18:CLASS 19:CLASS 22:CLASS 23:CLASS 24:CLASS 25:CLASS 26:CLASS 27:CLASS
- 28:CLASS 29:CLASS

L8 STRUCTURE UPLOADED

=> s 18

SAMPLE SEARCH INITIATED 10:37:38 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED -

100.0% PROCESSED

0 ITERATIONS

0 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: PROJECTED ANSWERS:

0 TO 0

1.9

0 SEA SSS SAM L8

=> d 18

L8 HAS NO ANSWERS

L8 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation.

=> s 18 sss full

FULL SEARCH INITIATED 10:38:03 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 1 TO ITERATE

SEARCH TIME: 00.00.01

1 ITERATIONS 0 ANSWERS

100.0% PROCESSED

L10 0 SEA SSS FUL L8

=> file stnguide

COST IN U.S. DOLLARS

SINCE FILE TOTAL

FULL ESTIMATED COST

ENTRY SESSION 374.64 374.86

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FILE CONTAINS CURRENT INFORMATION.

LAST RELOADED: Sep 25, 2009 (20090925/UP).

=> file hcaplus

COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.63 375.49

FULL ESTIMATED COST

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USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Aug 2009

 ${
m HCAplus}$ now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2009.

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=> s (lysophosphtidic acid)(4a)(inhib? or antag?)

0 LYSOPHOSPHTIDIC

4911805 ACID

0 LYSOPHOSPHTIDIC ACID

(LYSOPHOSPHTIDIC(W)ACID)

2188443 INHIB?

334665 ANTAG?

0 (LYSOPHOSPHTIDIC ACID) (4A) (INHIB? OR ANTAG?)

=> s phosphate or phrophosphate or phosphoric

637370 PHOSPHATE

4 PHROPHOSPHATE

121867 PHOSPHORIC

L12 717126 PHOSPHATE OR PHROPHOSPHATE OR PHOSPHORIC

=> s cholesterol or hypercholesterol? or hyperlipid? or atehrosclerosis or neointima

203298 CHOLESTEROL

20115 HYPERCHOLESTEROL?

18730 HYPERLIPID?

0 ATEHROSCLEROSIS

2232 NEOINTIMA

L13 220274 CHOLESTEROL OR HYPERCHOLESTEROL? OR HYPERLIPID? OR ATEHROSCLEROS
TS OR NEOTNTIMA

=> s 111 and 112

L14 0 L11 AND L12

=> s 111 and 113

0 L11 AND L13 1.15

=> s 112 and 113

6976 L12 AND L13 L16

=> file stnguide

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION

FULL ESTIMATED COST 2.85 378.34

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FILE CONTAINS CURRENT INFORMATION. LAST RELOADED: Sep 25, 2009 (20090925/UP).

=> file hcaplus COST IN U.S. DOLLARS

SINCE FILE TOTAL
ENTRY SESSION
207 378.41 FULL ESTIMATED COST

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FILE COVERS 1907 - 1 Oct 2009 VOL 151 ISS 14 FILE LAST UPDATED: 30 Sep 2009 (20090930/ED) REVISED CLASS FIELDS (/NCL) LAST RELOADED: Aug 2009 USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Aug 2009

HCAplus now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2009.

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The ALL, BIB, MAX, and STD display formats in the CA/CAplus family of databases have been updated to include new citing references

```
information. This enhancement may impact record import into
database management software. For additional information, refer
to NEWS 9.
=> s (lysophosphatidic acid)(4a)(inhib? or antag?)
          3893 LYSOPHOSPHATIDIC
       4911805 ACID
          3121 LYSOPHOSPHATIDIC ACID
                (LYSOPHOSPHATIDIC(W)ACID)
       2188443 INHIB?
        334665 ANTAG?
L17
           314 (LYSOPHOSPHATIDIC ACID) (4A) (INHIB? OR ANTAG?)
=> s phosphate or pyrophosphate or phosphoric
        637370 PHOSPHATE
         44113 PYROPHOSPHATE
        121867 PHOSPHORIC
L18
        744163 PHOSPHATE OR PYROPHOSPHATE OR PHOSPHORIC
=> s cholesterol or hypercholesterol? or hyperlipid? or atehrosclerosis or neointima
        203298 CHOLESTEROL
         20115 HYPERCHOLESTEROL?
         18730 HYPERLIPID?
            0 ATEHROSCLEROSIS
          2232 NEOINTIMA
L19
        220274 CHOLESTEROL OR HYPERCHOLESTEROL? OR HYPERLIPID? OR ATEHROSCLEROS
               IS OR NEOINTIMA
=> s 117 and 118
1.20
          62 L17 AND L18
=> s 117 and 119
L21
           3 L17 AND L19
=> s 118 and 119
         7389 L18 AND L19
=> file stnguide
COST IN U.S. DOLLARS
                                                 SINCE FILE
                                                                 TOTAL.
                                                      ENTRY
                                                              SESSION
FULL ESTIMATED COST
                                                       2.85
                                                               381.26
FILE 'STNGUIDE' ENTERED AT 10:44:25 ON 01 OCT 2009
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FILE CONTAINS CURRENT INFORMATION.
LAST RELOADED: Sep 25, 2009 (20090925/UP).
=> d 121 1-3 ti abs bib
YOU HAVE REQUESTED DATA FROM FILE 'HCAPLUS' - CONTINUE? (Y)/N:v
```

- L21 ANSWER 1 OF 3 HCAPLUS COPYRIGHT 2009 ACS on STN
- Anti-atherosclerotic molecules targeting oxidative stress and inflammation A review. The accumulation of lipids within arteries remains to be the AR initial impulse for the pathogenesis of atherosclerosis; however, both inflammation and oxidative stress are considered to play a critical role in this process. Several lipid lowering drugs are used as the first line therapy in atherosclerosis; however, different agents have been found to exhibit beneficial effects which are independent of their lipid lowering activity. Both statins and fibrates have been reported to exert anti-inflammatory and anti-oxidative effects in addition to their anti-atherosclerotic actions. Furthermore, anti-hypertensive, anti-diabetic and anti-platelet drugs, which reduce oxidative stress and inflammation, have been shown to attenuate atherosclerosis. In addition, novel substances such as HDL-related agents, cyclopentenone prostaglandins, lipoprotein-associated phospholipase A2 inhibitors, 5-lipoxygenase pathway inhibitors, acyl CoA: cholesterol acyltransferase inhibitors, analogs of probucol and lysophosphatidic acid antagonists have been developed for the treatment of atherosclerosis as a consequence of their actions on oxidative stress and inflammation. The present article reviews the involvement of inflammation and oxidative stress in the pathogenesis of atherosclerosis and focuses on the mechanisms of some clin, used as
- and anti-oxidative properties.
 AN 2009:1022565 HCAPLUS <<LOGINID::20091001>>
- DN 151:235384
- TI Anti-atherosclerotic molecules targeting oxidative stress and inflammation

well as potential anti-atherosclerotic substances with anti-inflammatory

- AU Adameova, A.; Xu, Y. J.; Duhamel, T. A.; Tappia, P. S.; Shan, L.; Dhalla, N. S.
- CS Institute of Cardiovascular Sciences, St. Boniface General Hospital Research Centre, Faculty of Medicine, University of Manitoba, Winnipeg, Can.
- SO Current Pharmaceutical Design (2009), 15(27), 3094-3107 CODEN: CPDEFP; ISSN: 1381-6128
- PB Bentham Science Publishers Ltd.
- DT Journal: General Review
- LA English
- RE.CNT 159 THERE ARE 159 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L21 ANSWER 2 OF 3 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Treatment of viral infections by modulation of host cell metabolic pathways
- AB Alterations of certain metabolite concns. and fluxes that occur in response to viral infection are described. Host cell enzymes in the involved metabolic pathways are selected as targets for intervention; i.e., to restore metabolic flux to disadvantage viral replication, or to further derange metabolic flux resulting in "suicide" of viral-infected cells (but not uninfected cells) in order to limit viral propagation. While any of the enzymes in the relevant metabolic pathway can be selected, pivotal enzymes at key control points in these metabolic pathways are preferred as candidate antiviral drug targets. Inhibitors of these enzymes are used to reverse, or redirect, the effects of the viral infection. Drug candidates are tested for antiviral activity using screening assays in vitro and host cells, as well as in animal models. Animal models are then used to test efficacy of candidate compds. in preventing and treating viral infections. The antiviral activity of enzyme inhibitors is demonstrated.
- AN 2009:198413 HCAPLUS <<LOGINID::20091001>>
- DN 150:252581

```
Treatment of viral infections by modulation of host cell metabolic
     pathways
TN
     Shenk, Thomas; Rabinowitz, Joshua D.; Munger, Josh; Bennett, Bryson
PΑ
    The Trustees of Princeton University, USA
SO
    PCT Int. Appl., 339pp.
     CODEN: PIXXD2
     Patent
LA
     English
FAN.CNT 1
                     KIND DATE APPLICATION NO. DATE
    PATENT NO.
    WO 2009023059 A2 20090219 WO 2008-US6959 20080602
        W: AE, AG, AL, AM, AO, AT, AU, AZ, BA, BB, BG, BH, BR, BW, BY, BZ,
             CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES,
             FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE,
             KG, KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD,
             ME, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH,
             PL, PT, RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM,
             TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW
         RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HR, HU,
             IE, IS, IT, LT, LU, LV, MC, MT, NL, NO, PL, PT, RO, SE, SI, SK,
             TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW,
             AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
US 20090239830 A1 20090924
PRAI US 2007-932769P P 20070601
US 2008-33243P P 20080303
                                           US 2008-156517
ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
   MARPAT 150:252581
L21 ANSWER 3 OF 3 HCAPLUS COPYRIGHT 2009 ACS on STN
    Lysophosphatidic acid analogs and inhibition
ΤI
     of neointima formation
AB
    The phospholipid growth factor lysophosphatidic acids (LPAs) containing
     unsatd. fatty acids (18:1, 18:2 and 20:4) and fatty alcs. containing
     hydrocarbon chains with more than 4 carbons were capable of inducing a
     rapid formation of neointima, an initial step in the development
     of atherosclerotic plaque. LPAs with saturated fatty acids did not induce
     neointima formation. A Peroxisome Proliferator-Activated
     Receptors gamma (PPARy)-specific agonist Rosiglitasone also induced
     a profound formation of neointima. GW9662, a selective and
     irreversible antagonist of PPARy, abolished LPA- and
     Rosiglitazone-induced neointima formation, indicating that
     LPA-induced neointima formation requires the activation of
     PPARy. These data suggest that LPA analogs that bind to but do not
     activate downstream signaling of PPARy or antagonists of PPARy
     that inhibit PPARy signaling would be useful in the prevention
     and/or treatment of neointima formation and atherosclerosis.
AN
     2004:857161 HCAPLUS <<LOGINID::20091001>>
DN
     141:343506
ΤI
     Lysophosphatidic acid analogs and inhibition
     of neointima formation
IN
     Tigvi, Gabor; Baker, Daniel L.; Zhang, Chunxiang
PA
SO
    U.S. Pat. Appl. Publ., 23 pp.
    CODEN: USXXCO
    Patent
LA English
FAN.CNT 1
     PATENT NO. KIND DATE APPLICATION NO. DATE
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PI	US	20040204383								US 2004-821739						20040409					
	AU	2004229467				A1		2004	1028		AU 2004-229467					20040409					
	AU	2004229467				B2	32 20070125														
	CA	2521189				A1		2004	1028		CA 2	004-	2521	20040409							
	WO	2004	0914	96		A2		2004	1028		WO 2	004-	US11	20040409							
	WO	2004						0011010													
		W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,			
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		DM.							MZ,												
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		TD, TG													00010100						
	EP	1613298				A2 20060111 CH, DE, DK, ES, FR,															
		R:																			
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PRAI		2003-462274P																			
	WO	2004		W		20040409															

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USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Aug 2009

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L23 32 L20 AND (PY<2003 OR AY<2003 OR PRY<2003)

=> d 123 1-32 ti abs bib hitstr

L23 ANSWER 1 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN

TI The phospholipids sphingosine-1-phosphate and lysophosphatidic acid prevent apoptosis in osteoblastic cells via a signaling pathway involving Gi proteins and phosphatidylinosito1-3 kinase

- AB The naturally occurring phospholipids lysophosphatidic acid (LPA) and sphingosine-1-phosphate (S1P) have recently emerged as bioactive compds. that exert mitogenic effects in many cell types, including osteoblasts. In the current study, we examined the ability of each of these compds. to influence osteoblast survival. Using terminal deoxynucleotidyl transferase-mediated deoxyuridine 5'-triphosphate nick-end labeling and DNA fragmentation assays, we found that both LPA and S1P dose-dependently inhibited (by at least 50% and 40%, resp.) the apoptosis induced by serum withdrawal in cultures of primary calvarial rat osteoblasts and SaOS-2 cells. The antiapoptotic effects were inhibited by pertussis toxin, wortmannin, and LY294002, implicating Gi proteins and phosphatidylinositol-3 kinase (PI-3 kinase) in the signaling pathway that mediates phospholipid-induced osteoblast survival. Specific inhibitors of p42/44 MAPK signaling did not block LPA- or S1P-induced osteoblast survival. LPA and S1P induced PI-3 kinase-dependent activation of p70 S6 kinase, but rapamycin, a specific inhibitor of p70 S6 kinase activation, did not prevent phospholipid-induced osteoblast survival. LPA and SIP also inhibited apoptosis in Swiss 3T3 fibroblastic cells in a Gi protein-dependent fashion. In fibroblastic cells, however, the antiapoptotic effects of SIP were sensitive to inhibition of both PI-3 kinase and p42/44 MAPK signaling, whereas those of LPA were partially abrogated by inhibitors of p42/44 MAPK signaling but not by PI-3 kinase inhibitors. These data demonstrate that LPA and SIP potently promote osteoblast survival in vitro, and that cell-type specificity exists in the antiapoptotic signaling pathways activated by phospholipids.
- AN 2002:923658 HCAPLUS <<LOGINID::20091001>>

DN 138:318116

- II The phospholipids sphingosine-1-phosphate and lysophosphatidic acid prevent apoptosis in osteoblastic cells via a signaling pathway involving Gi proteins and phosphatidylinositol-3 kinase
- AU Grey, Andrew; Chen, Qi; Callon, Karen; Xu, Xin; Reid, Ian R.; Cornish, Jill
- CS Department of Medicine, University of Auckland, Auckland, N. Z.
- Endocrinology (2002), 143(12), 4755-4763 CODEN: ENDOAO; ISSN: 0013-7227
- PB Endocrine Society
- DT Journal

LA English

OSC.G 37 THERE ARE 37 CAPLUS RECORDS THAT CITE THIS RECORD (37 CITINGS)

- L23 ANSWER 2 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- Comparative analysis of human and rat S1P5 (edg8): differential expression profiles and sensitivities to antagonists
- AB Five quanine nucleotide-binding protein-coupled receptors (S1P1-5) for the lysophospholipid mediator sphingosine 1-phosphate (S1P) have thus far been described. Whereas tissue distribution and functional properties of the human S1P1-4 genes are well characterized, only limited functional and expression data are available for S1P5, to date. Northern blot anal. indicated that human S1P5 (hS1P5) is an alternatively spliced gene, with a 5.4-kb transcript that is predominantly expressed in peripheral tissues, and a 2.4-kb transcript expressed in brain, spleen, and peripheral blood leukocytes. In contrast, rat S1P5 (rS1P5) was exclusively detected in brain and skin. Expression of hS1P5 and rS1P5 in mammalian CHO-K1 or HEK293 cells conferred onto the cells the ability to mobilize intracellular calcium as determined by a functional Fluorometric Imaging Plate Reader assay, when challenged with S1P and dihydro S1P, resp. Applying a lipid library with 200 bioactive lipids in a functional Fluorometric Imaging Plate Reader assay did not reveal addnl. agonists. However, both receptors exhibited differential sensitivity towards the S1P- and lysophosphatidic acid-receptor antagonist, suramin: rS1P5-mediated intracellular calcium

mobilization was partly inhibited by suramin (IC50: 5800 μM), whereas hS1P5 was completely antagonized (IC50: 130 µM). Both receptors were sensitive towards inhibition with the related drug (8,8'-(carbonylbis(imino-3,1-phenylene))bis(1,3,5naphthalenetrisulfonicacid))but ic50 values differed significantly (340 μM for hS1P5, 4000 μM for rS1P5). In addition, rS1P5 displayed antiproliferative effects in transfected CHO-K1 and HEK293 cells in

contrast to hS1P5. Taken together, our data imply that differences between hS1P5 and rS1P5 will be an important point to be considered in the development of selective receptor antagonists.

- 2002:701945 HCAPLUS <<LOGINID::20091001>> AN
- DN 138:317912
- TΙ Comparative analysis of human and rat S1P5 (edg8): differential expression profiles and sensitivities to antagonists
- AU Niedernberg, Anke; Scherer, Constanze R.; Busch, Andreas E.; Kostenis, Evi
- CS Disease group Cardiovascular, Frankfurt, 65926, Germany
- SO Biochemical Pharmacology (2002), 64(8), 1243-1250 CODEN: BCPCA6: ISSN: 0006-2952
- PB Elsevier Science Inc.
- DT Journal
- LA. English
- OSC. G 1.0
- THERE ARE 10 CAPLUS RECORDS THAT CITE THIS RECORD (10 CITINGS) RE.CNT 42 THERE ARE 42 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L23 ANSWER 3 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- Modification of the antitumor effect of doxorubicin by phosphorylated ΤI retinoids conjugated to α-fetoprotein
- The influence of new retinoids representing all-trans-retinoic acid (RA) amides with O-phosphates of ethanolamine, L-serine, L-threonine (Thr), and L-tyrosine conjugated to a-fetoprotein on the antitumor action of the antibiotic doxorubicin was evaluated with regard to inoculated Ehrlich ascitic carcinoma. The synthesized compds. are structurally analogous to N-palmitoy1-0-phospho-L-serine (NP-Ser-PA) and N-palmitoy1-O-phospho-L-tyrosine (NP-Tyr-PA), which are antagonists of the receptors of lysophosphatidic acid. The complex of doxorubicin with NR-Thr-PA in the presence

of $\alpha\text{-FP}$ displayed a lower antitumor activity compared to that of free doxorubicin, in which the Ehrlich carcinoma cell growth was inhibited by 37%. The RA complex containing only NR-Tyr-PA produced a more evident cytotoxic action upon the tumor compared to both the complex and free doxorubicin. An optimum composition was offered by the doxorubicin complex with NR-Tyr-PA and $\alpha\text{-FP}$ produced a pos. influence on the antitumor properties of this complex. The absence of a reliable effect for the mixture of doxorubicin with NR-Tyr-PA is attributed by the formation of an ion complex between the components, related to the interaction between the amino group of daunosamine and the phosphate moiety of NR-Tyr-PA.

- AN 2002:628766 HCAPLUS <<LOGINID::20091001>>
- DN 138:147338
- TI Modification of the antitumor effect of doxorubicin by phosphorylated retinoids conjugated to α -fetoprotein
- AU Arsenov, D. V.; Babitskaya, S. V.; Vashkevich, I. I.; Dad'kov, I. D.; Kisel', M. A.; Kuz'mitskii, B. B.; Strel'chenok, O. A.
- CS Institute of Bioorganic Chemistry, National Academy of Sciences of Belarus, Minsk, Belarus
- SO Pharmaceutical Chemistry Journal (Translation of Khimiko-Farmatsevticheskii Zhurnal) (2001), 35(12), 657-660 CODEN: PCJOAU; ISSN: 0991-150X
- PB Kluwer Academic/Consultants Bureau
- DT Journal
- LA English
- RE.CNT 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L23 ANSWER 4 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Molecular modeling of lysophosphatidic acid receptor antagonists
- AB Lysophosphatidic acid (LPA) elicits a variety of responses including mitogenesis, cytoskeletal changes, activation of Ca2+ transients, and effects on apoptosis. These responses are elicited via LPA1/SDG2, LPA2/SDG4, and LPA3/SDGG G protein-coupled receptors. These receptors are members of the endothelial differentiation gene family. In order to understand the physiol. significance of LPA highly selective antagonists are necessary. Recently, dioctyl glycerol pyrophosphate (DGPP) and fatty alkyl phosphate (FAP) were shown to be potent and selective antagonists towards LPA3 receptor. We have docked DGPP and FAP in our LPA1, LPA2, and LPA3 receptor models and our docked energies agree with the observed trend in inhibition consts. (Ki). The docked positions of the antagonist relative to the agonist overlap in the position of the polar head group, but diverge in the favored position of the hydrophobic tail(s).
- AN 2002:617933 HCAPLUS <<LOGINID::20091001>>
- TI Molecular modeling of lysophosphatidic acid receptor antagonists
- AU Sardar, Vineet M.; Virag, Tamas; Fischer, David J.; Elrod, Don; Bautista, Debra L.; Wang, De-an; Nusser, Nora; Yokoyama, Kazuaki; Baker, Daniel L.; Miller, Duane D.; Tigyi, Gabor; Parrill, Abby L.
- CS Department of Chemistry and Computational Research on Materials Institute, University of Memphis, Memphis, TN, 38152-6060, USA
- SO Abstracts of Papers, 224th ACS National Meeting, Boston, MA, United States, August 18-22, 2002 (2002), MEDI-079 Publisher: American Chemical Society, Washington, D. C. CODEN: 69CZPZ
- DT Conference; Meeting Abstract
- LA English
- L23 ANSWER 5 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN

- TI Lysophosphatidic acid inhibition of the accumulation of Pseudomonas aeruginosa PAOl alginate, pyoverdin, elastase and LasA
- AB The pathogenesis of Pseudomonas aeruginosa is at least partially attributable to its ability to synthesize and secrete the siderophore pyoverdin and the two zinc metalloproteases elastase and LasA, and its ability to form biofilms in which bacterial cells are embedded in an alginate matrix. In the present study, a lycophospholipid, 1-palmitoy1-2-hydroxy-sn-glycero-3-phosphate [also called monopalmitoy]phosphatidic acid (MPPA)], which accumulates in inflammatory exudates, was shown to inhibit the extracellular accumulation of P. aeruginosa PAOI alginate, elastase, LasA protease and the siderophore pyoverdin. MPPA also inhibited biofilm formation. The inhibitory effects of MPPA occur independently of rpoS expression and without affecting the accumulation of the autoinducers N-(3-oxododecanoyl) homoserine lactone and N-butyryl-L-homoserine lactone, and may be due, at least in part, to the ability of MPPA to divalent cations.
- AN 2002:484327 HCAPLUS <<LOGINID::20091001>>
- DN 138:86340
- TI Lysophosphatidic acid inhibition of the
- accumulation of Pseudomonas aeruginosa PAO1 alginate, pyoverdin, elastase and LasA AU Laux, David C.; Corson, Jov M.; Givskov, Michael; Hentzer, Morten; Moller,
- Annette; Wosencroft, Kathleen A.; Olson, Joan C.; Krogfelt, Karen A.; Goldberg, Joanna B.; Cohen, Paul S.
- CS Department of Cell and Molecular Biology, University of Rhode Island, Kingston, RI, 02881, USA
- SO Microbiology (Reading, United Kingdom) (2002), 148(6), 1709-1723 CODEN: MROBEO; ISSN: 1350-0872
- PB Society for General Microbiology
- DT Journal
- LA English
- OSC.G 6 THERE ARE 6 CAPLUS RECORDS THAT CITE THIS RECORD (6 CITINGS)
 RE.CNT 68 THERE ARE 68 CITED REFERENCES AVAILABLE FOR THIS RECORD
 ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L23 ANSWER 6 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Molecular basis for lysophosphatidic acid receptor antagonist selectivity
- AB A review. Recent characterization of lysophosphatidic acid (LPA) receptors has made possible studies elucidating the structure-activity relationships (SAR) for agonist activity at individual receptors. Addnl., the availability of these receptors has allowed the identification of antagonists of LPA-induced effects. Two receptor-subtype selective LPA receptor antagonists, one selective for the LPA1/EDG2 receptor (a benzyl-4-oxypenzyl N-acyl ethanolamide phosphate, NAEPA, derivative) and the other selective for the LPA3/EDG7 receptor (diacylglycerol pyrophosphate, DGPP, 8:0), have recently been reported. The receptor SAR for both agonists and antagonists are reviewed, and the mol. basis for the difference between agonism and antagonism as well as for receptor-subtype antagonist selectivity identified by mol. modeling is described. The implications of the newly available receptor-subtype selective antagonists are also discussed.
- AN 2002:459276 HCAPLUS <<LOGINID::20091001>>
- DN 138:86877
 - Molecular basis for lysophosphatidic acid receptor
 - antagonist selectivity
- AU Sardar, Vineet M.; Bautista, Debra L.; Fischer, David J.; Yokoyama, Kazuaki; Nusser, Nora; Virag, Tamas; Wang, De-an; Baker, Daniel L.; Tigyi, Gabor; Parrill, Abby L.
- CS Department of Chemistry and Computational Research on Materials Institute,

The University of Memphis, Memphis, TN, 38152-6060, USA

- Biochimica et Biophysica Acta, Molecular and Cell Biology of Lipids (SO 2002), 1582(1-3), 309-317 CODEN: BBMLFG; ISSN: 1388-1981
- PB Elsevier B.V.
- DT Journal: General Review
- LA English
- osc.g 52 THERE ARE 52 CAPLUS RECORDS THAT CITE THIS RECORD (53 CITINGS)
- RE.CNT 32 THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD
 - ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L23 ANSWER 7 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- ΤI Albumin stimulates lysophosphatidic acid acyltransferase activity in T-lymphocyte membranes
- AB Phosphatidic acid (PtdOH) and lysophosphatidic acid (lysoPtdOH) have been shown to enhance T-lymphocyte function. However, the FA preference and influence of acyl-CoA binding proteins on lysoPtdOH and PtdOH biosynthesis are not known. Therefore, we determined glycerol-3-phosphate acyltransferase (GPAT) and lysophosphatidic acid acyltransferase (LAT) activity in rat T-lymphocyte and liver membrane prepns. in the presence of palmitoy1-CoA and oleoy1-CoA with or without BSA. We found two different properties of GPAT and LAT in whole T-lymphocyte membrane prepns. relative to liver. First, T-lymphocyte basal GPAT and LAT activities were similar, whereas in liver membranes LAT activity was 10-fold higher than GPAT. Second, T-lymphocyte LAT, but not GPAT, activity was inducible (fivefold) by the addition of albumin in the presence of palmitoyl-CoA but not oleoyl-CoA. In contrast, albumin stimulated GPAT, but not LAT, activity in liver membranes in the presence of palmitoyl-CoA. These results show, for the first time, that T-lymphocyte LAT activity can be increased by the presence of an acyl-CoA binding protein, which may indicate a new important control mechanism for regulating intracellular lysoPtdOH and PtdOH levels in T-lymphocytes.
 - 2002:396235 HCAPLUS <<LOGINID::20091001>>
- AN 137:198675 DN
- TI Albumin stimulates lysophosphatidic acid acyltransferase activity in T-lymphocyte membranes
- ΑU Jolly, Christopher A.; Kannan, Latha
- CS Division of Nutritional Sciences and Institute for Cellular and Molecular Biology, The University of Texas at Austin, Austin, TX, 78712, USA
- so Lipids (2002), 37(5), 475-480
 - CODEN: LPDSAP; ISSN: 0024-4201
- PB AOCS Press
- DT Journal LA English
- OSC.G 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD (4 CITINGS)
- RE.CNT 25 THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L23 ANSWER 8 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- TΙ Noradrenaline release-inhibiting receptors on PC12 cells devoid of α2- and CB1 receptors: similarities to presynaptic imidazoline and edg receptors
- on rat pheochromocytoma PC12 cells. Veratridine-evoked [3H]noradrenaline release from PC12 cells was inhibited by micromolar concns. of the imidazoline and quanidine derivs. cirazoline, clonidine, aganodine, 1,3-di(2-toly1)guanidine, BDF6143 and agmatine, and of the cannabinoid receptor agonist WIN55,212-2 (R(+)-[2,3-dihydro-5-methyl-3-[(morpholinyl)methyl]pyrrolo-[1,2,3-de]-1,4-benzoxazin-yl](1naphthalenyl) methanone mesylate), but not by noradrenaline. The inhibitory effect of clonidine was antagonized by micromolar concns. of

The aim of the present study was to classify release-inhibiting receptors

rauwolscine and SR141716A (N-[piperidin-1-y1]-5-[4-chloropheny1]-1-[2,4dichlorophenyl | -4-methyl-1H-pyrazole-3-carboxamide). The potencies of the agonists and antagonists were compatible with an action at previously characterized presynaptic imidazoline receptors. 1-Oleoyl-lysophosphatidic acid, but not sphingosine-1-phosphate, produced an inhibition of release that was antagonized by 30 µM rauwolscine, 1 µM SR141716A and 10 µM LY320135 as well as by pretreatment of the cells with 100 µM clonidine for 72 h. Polymerase chain reaction (PCR) expts. on cDNA from PC12 mRNA suggest mRNA expression of lysophospholipid receptors encoded by the genes edg2, edg3, edg5 and edg7, but not of receptors encoded by edg1, edg4, edg6 and edg8, and not of a2A-nd CB1 receptors. In conclusion, PC12 cells are not endowed with a2-adrenoceptors and CB1 cannabinoid receptors, but with an inhibitory receptor recognizing imidazolines, guanidines and WIN55,212-2 similar to that on sympathetic nerves. The PCR results and the ability of 1-oleoyl-LPA to mimic these drugs (also with respect to their susceptibility to antagonists) suggest that the release-inhibiting receptor may be an edg-encoded lysophospholipid receptor. 2001:883810 HCAPLUS <<LOGINID::20091001>>

AN 2001:883810 DN 136:319263

TI Noradrenaline release-inhibiting receptors on PC12 cells devoid of $\alpha 2-$ and CBI receptors: similarities to presynaptic imidazoline and edd receptors

AU Molderings, G. J.; Bonisch, H.; Hammermann, R.; Gothert, M.; Bruss, M. CS Institute of Pharmacology and Toxicology, University of Bonn, Bonn, 53113, Germany

SO Neurochemistry International (2002), 40(2), 157-167

CODEN: NEUIDS; ISSN: 0197-0186

PB Elsevier Science Ltd. DT Journal

DT Journal LA English

OSC.G 21 THERE ARE 21 CAPLUS RECORDS THAT CITE THIS RECORD (21 CITINGS)
RE.CNT 26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L23 ANSWER 9 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN

TI Short-chain phosphatidates are subtype-selective antagonists of lysophosphatidic acid receptors

AB Lysophosphatidic acid (LPA) and sphingosine-1-phosphate (SIP) are members of the phospholipid growth factor family. A major limitation in the field to date has been a lack of receptor subtype-specific agonists and antagonists. Here, we report that dioctylglycerol pyrophosphate and dioctylphosphatidic acid are selective antagonists of the LPA1 and LPA3 receptors, but prefer LPA3 by an order of

antagonists of the LPAI and LPAI receptors, but prefer LPAI by an order of magnitude. Neither mol. had an agonistic or antagonistic effect on LPA2 receptor. Consistent with this receptor subtype selectivity, dioctylglycerol pyrophosphate inhibited cellular responses to LPA in NiH3T3 fibroblasts, HEY ovarian cancer cells, PC12 pheochromocytoma cells, and Xenopus laevis occytes. Responses elicited by SIP in these cell lines that endogenously express SIP1, SIP2, SIP3, and SIP5 receptors were unaffected by dioctylglycerol pyrophosphate. Responses evoked by the G protein-coupled receptor liquads acetylcholine, serotonin, ATP, and thrombin receptor-activating pertide were similarly unaffected,

subtype-specific lysophosphatidate antagonists.

AN 2001:723801 HCAPLUS <<LOGINID::20091001>> DN 136:96559

TI Short-chain phosphatidates are subtype-selective antagonists of lysophosphatidic acid receptors

suggesting that the short-chain phosphatidates are receptor

AU Fischer, David J.; Nusser, Nora; Virag, Tamas; Yokoyama, Kazuaki; Wang, De-An; Baker, Daniel L.; Bautista, Debra; Parrill, Abby L.; Tigyi, Gabor

- CS Department of Physiology, University of Tennessee Health Science Center, Memphis, TN, USA
- SO Molecular Pharmacology (2001), 60(4), 776-784 CODEN: MOPMA3; ISSN: 0026-895X
- PB American Society for Pharmacology and Experimental Therapeutics
- DT Journal
- LA English
- OSC.G 89 THERE ARE 89 CAPLUS RECORDS THAT CITE THIS RECORD (89 CITINGS)
- RE.CNT 36 THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L23 ANSWER 10 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Synthesis of lysophosphatidic acid receptor agonists and antagonists and their use for cancer inhibition, wound healing, and enhancement of cell proliferation
- AB The present invention relates to lysophosphatidic acid (LPA) analogs and cyclic derivs. of the analogs as well as pharmaceutical compns. which include those compds. Also disclosed are methods of using such compds., which have activity as agonists or as antagonists of LPA receptors; such methods including inhibiting LPA activity on an LPA receptor, modulating LPA receptor activity, treating cancer, enhancing cell proliferation, and treating a wound. Thus, 2-amino-3-oxo-3-(tetradecylamino)propyl dihydrogen phosphate (1),
 - 2-(acetylamino)-3-oxo-3-(tetradecylamino)propyl dihydrogen phosphate (II), and 1,2-(3-octadecyloxypropane)-bis(dihydrogen phosphate) (III) were synthesized. The cytotoxicity of these
- phosphate) (III) were synthesized. The cytotoxicity of these compds. on prostate cancer cell lines was determined The IC50's observed were 0.7
- \pm 0.1 for I on PC-3 cells, 0.7 \pm 0.1 for II on DU145 cells, and 3.1 \pm 3.2 for III on LNCaP cells. Addnl., phosphoric acid monododecyl ester (IV) was prepared and screened in Xenopus oocytes (which produce the PS224 receptor) and in recombinant RH7777 cells producing Edg-2, Edg-4, and Edg-7 receptors. In Xenopus IV inhibited LPA-induced chloride currents with an IC50 value of about 8.1 nM. In Edg-2 and Edg-4-expressing RH7777 cells IV visingificantly inhibited the Ca2+ responses while in Edg-7-expressing cells this compound increased the Ca2+ responses.
- AN 2001:713600 HCAPLUS <<LOGINID::20091001>>
- DN 135:267219
- TI Synthesis of lysophosphatidic acid receptor agonists and antagonists and their use for cancer inhibition, wound healing, and enhancement of cell proliferation
- IN Miller, Duane D.; Tigyi, Gabor; Dalton, James T.; Sardar, Vineet M.; Elrod, Don B.; Xu, Huiping; Baker, Daniel L.; Wang, Dean; Liliom, Karoly; Fischer, David J.; Viraq, Tamae; Nusser, Nora
- PA University of Tennessee Research Corporation, USA
- SO PCT Int. Appl., 140 pp.
- CODEN: PIXXD2
- DT Patent
- LA English FAN.CNT 3
- PATENT NO. KIND DATE APPLICATION NO. DATE

 PI WO 2001071022 A2 20010927 WO 2001-US8729 20010319 <--
 - N: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
 - SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW

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RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
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                         A1
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     AU 2001049263
                         Α
                                20011003
                                           AU 2001-49263
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     EP 1263752
                         A2
                               20021211
                                           EP 2001-922465
                                                                   20010319 <---
     EP 1263752
                               20071205
                         B1
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
     JP 2004506604
                         Т
                               20040304
                                           JP 2001-569403
                                                                   20010319 <--
     AT 380187
                         Т
                               20071215
                                           AT 2001-922465
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     EP 1918287
                         A2
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     EP 1918287
                         A3
                               20080820
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     ES 2298227
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                               20080516
                                           ES 2001-922465
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     KR 874392
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     AU 2007202615
                        A1
                               20070628
                                           AU 2007-202615
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                               20000317 <--
PRAI US 2000-190370P
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                         A3
                               20010319 <--
     EP 2001-922465
                               20010319
                         A3
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     WO 2001-US8729
                         W
                               20010319
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    MARPAT 135:267219
OSC.G
             THERE ARE 6 CAPLUS RECORDS THAT CITE THIS RECORD (7 CITINGS)
L23 ANSWER 11 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
    Synthesis of N-arachidonyl-O-phospho-L-serine and of
     N-arachidonov1-O-phospho-L-tyrosine as nonsaturated lysophosphatic acid
    receptor antagonists
AR
    A method for preparing unsatd. antagonists of
    lysophosphatidic acid receptor was developed. The
     synthesis of both N-arachidonyl-O-phospho-L-serine (I) and
     N-arachidonyl-O-phospho-L-tyrosine is described. I was prepared by the
     reaction of L-serine with arachidonic acid in the presence of Et3N
     followed by the reaction of the resulting N-arachidonyl-L-serine with
     β-cyanoethyl phosphate.
AN
    2001:601416 HCAPLUS <<LOGINID::20091001>>
DN
    136:330465
ΤI
    Synthesis of N-arachidonyl-O-phospho-L-serine and of
    N-arachidonov1-O-phospho-L-tyrosine as nonsaturated lysophosphatic acid
    receptor antagonists
AU
    Arsenov, D. V.; Kisel, M. A.; Strel'chenok, O. A.
CS
    Inst. Bioorg. Khim., NAN Belarusi, Belarus
SO
    Doklady Natsional'noi Akademii Nauk Belarusi (2001), 45(3),
     71 - 73
     CODEN: DNABFW; ISSN: 1561-8323
PB
     Belaruskava Navuka
DT
    Journal
LA
    Russian
OSC.G 1
             THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)
L23 ANSWER 12 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
     Lysophosphatidic acid inhibits Ca2+
     signaling in response to epidermal growth factor receptor stimulation in
     human astrocytoma cells by a mechanism involving phospholipase Cy
     and a Gai protein
AB
    The effect of the lysophospholipid mediators lysophosphatidic acid (LPA)
     and sphingosine 1-phosphate and the polypeptide growth factor
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epidermal growth factor (EGF) on the human astrocytoma cell line 1321N1 was assessed. These agonists produced a rapid and transient increase of the intracellular Ca2+ concentration When LPA was perfused before addition of EGF,

the EGF-dependent Ca2+ transient was abrogated, whereas this was not observed when EGF preceded LPA addition This inhibitory effect was not found for other EGF-mediated responses, e.g., activation of the mitogen-activated protein kinase cascade and cell proliferation, thus pointing to the existence of cross-talk between LPA and EGF for only a branch of EGF-induced responses. As 1321N1 cells expressed mRNA encoding the LPA receptors endothelial differentiation gene (Edg)-2, Edg-4, and Edg-7 and as sphingosine 1-phosphate did not interfere with LPA signaling, Edg-2, Edg-4, and/or Edg-7 could be considered as the LPA receptors mediating the aforementioned cross-talk. Attempts to address the biochem. mechanism involved in the cross-talk between the receptors were conducted by the immunopptn. approach using antibodies reacting with the EGF receptor (EGFR), phosphotyrosine, phospholipase Cy (PLCy)-1, and Gai protein. LPA was found to induce coupling of PLCy-1 to the EGFR by a mechanism involving a Gai protein, in the absence of tyrosine phosphorylation of both PLCy and the EGFR. These data show a cross-talk between LPA and EGF limited to a branch of EGFR-mediated signaling, which may be explained by a LPA-induced, Gai-protein-mediated translocation of PLCy-1 to EGFR in the absence of detectable tyrosine phosphorylation of both proteins.

AN 2000:675655 HCAPLUS <<LOGINID::20091001>> DN 133:291530

TI Lysophosphatidic acid inhibits Ca2+

signaling in response to epidermal growth factor receptor stimulation in human astrocytoma cells by a mechanism involving phospholipase Cy and a Gai protein

Hernandez, Marita; Barrero, Maria Jose; Crespo, Mariano Sanchez; Nieto, AU Maria Luisa

CS Instituto de Biologia y Genetica Molecular, CSIC-Universidad de Valladolid, Valladolid, 47005, Spain

SO Journal of Neurochemistry (2000), 75(4), 1575-1582

CODEN: JONRA9; ISSN: 0022-3042 Lippincott Williams & Wilkins

PB

DT Journal LA English

OSC.G 20 THERE ARE 20 CAPLUS RECORDS THAT CITE THIS RECORD (20 CITINGS) RE.CNT 44 THERE ARE 44 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L23 ANSWER 13 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN

Lysophosphatidic acid and sphingosine 1-phosphate stimulate TI

endothelial cell wound healing AB Lysophosphatidic acid (LPA) and sphingosine 1-phosphate (S1P) are potent lipid growth factors with similar abilities to stimulate cytoskeleton-based cellular functions. Their effects are mediated by a subfamily of G protein-coupled receptors (GPCRs) encoded by endothelial differentiation genes (edgs). The authors hypothesize that large quantities of LPA and S1P generated by activated platelets may influence endothelial cell functions. Using an in vitro wound healing assay, the authors observed that LPA and S1P stimulated closure of wounded monolayers of human umbilical vein endothelial cells and adult bovine aortic endothelial cells, which express LPA receptor Edg2, and S1P receptors Edg1 and Edg3. The two major components of wound healing, cell migration and proliferation, were stimulated individually by both lipids. LPA and S1P also stimulated intracellular Ca2+ mobilization and mitogen-activated protein kinase (MAPK) phosphorylation. Pertussis toxin partially blocked the effects of both lipids on endothelial cell migration, MAPK phosphorylation, and Ca2+ mobilization, implicating Gi/o-coupled Edg receptor signaling in endothelial cells. LPA and S1P did not cross-desensitize each other in Ca2+ responses, suggesting involvement of distinct receptors. Thus LPA and S1P affect endothelial cell functions

through signaling pathways activated by distinct GPCRs and may contribute to the healing of wounded vasculatures.

- AN 2000:189961 HCAPLUS <<LOGINID::20091001>>
- DN 132:320462
- TI Lysophosphatidic acid and sphingosine 1-phosphate stimulate endothelial cell wound healing
- AU Lee, Hsinyu; Goetzl, Edward J.; An, Songzhu
- CS Department of Medicine, University of California Medical Center, San Francisco, CA, 94143-0711, USA
- SO American Journal of Physiology (2000), 278(3, Pt. 1), C612-C618 CODEN: AJPHAP; ISSN: 0002-9513
- PB American Physiological Society
- DT Journal
- LA English
- OSC.G 122 THERE ARE 122 CAPLUS RECORDS THAT CITE THIS RECORD (122 CITINGS)
 RE.CNT 32 THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD
 - ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L23 ANSWER 14 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Methods using a lysophosphatidic acid receptor agonist for promoting survival of myelin-producing cells
- AB The invention is in the field of neurobiol., and relates particularly to methods useful for enhancing the survival of myelin producing cells, in particular Schwann cells and oligodendrocytes, and thereby to treating diseases of the nervous system involving loss of myelination or aberrant myelination. The methodol. of the invention uses a survival-promoting amount of an lysophosphatidic acid (LPA) receptor agonist, e.g. LPA.
- AN 2000:133529 HCAPLUS <<LOGINID::20091001>>
- DN 132:175856
- TI Methods using a lysophosphatidic acid receptor agonist for promoting survival of myelin-producing cells
- IN Chun, Jerold J. M.; Weiner, Joshua A.; Wickens, Philip L.; Begleiter, Leath E.
- PA The Regents of the University of California, USA; Allelix Biopharmaceuticals Inc.
- SO PCT Int. Appl., 37 pp.
- CODEN: PIXXD2
- DT Patent

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	WO 2000009139					A3 20000518														
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OSC.G 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD (3 CITINGS)

- L23 ANSWER 15 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- Enhancement of the Migration of Metastatic Human Breast Cancer Cells by Phosphatidic Acid
- Phosphatidic acid (PA), lysophosphatidic acid (LPA), and sphingosine 1phosphate (SPP) are naturally occurring phospholipids which induce a variety of effects as extracellular messengers. In this study, we compared the effects of these phospholipid signaling mols. on the migration of invasive and noninvasive breast cancer cell lines, an index of the metastatic potential of these cells. As previously demonstrated, invasive MDA-MB-231 breast cancer cells exhibited increased constitutive (nonstimulated) migration in comparison to poorly invasive MCF-7 cells. Phosphatidic acid employed at nanomolar concns. markedly potentiated migration of the invasive cells but had no effect on migration of either the noninvasive MCF-7 cells or nonneoplastic human epithelial cells. Lysophosphatidic acid and sphingosine 1phosphate inhibited both the directed (chemotactic) and random (chemokinetic) migration of MDA-MB-231 cells. Expts. were undertaken to characterize the signaling pathway involved in constitutive and PA-stimulated migration of MDA-MB-231 cells. The tyrosine kinase inhibitors staurosporine and genistein inhibited constitutive and PA-induced migration in a dose-dependent manner, consistent with a role for tyrosine phosphorylation in the migratory response. In addition, the phosphatidylinositol (PI) 3' kinase inhibitors wortmannin and LY294002

strongly inhibited both the constitutive and PA-stimulated migration of the invasive breast cancer cells, indicating that PI-3' kinase plays an important role in the metastatic migration of breast cancer cells. Finally, PA-induced migration of MDA-MB-231 was markedly attenuated by pretreatment of cells with Clostridium difficile Toxin B, pertussis toxin and suramin, implying a role for a Gi receptor-dependent process involving activation of the small GTP-binding protein Rho. Since an enhanced ability to migrate heightens the metastatic potential of cells within solid tumors, our results suggest that the metastatic capabilities of breast cancer cells may be enhanced by a receptor-driven cellular process initiated by phosphatidic acid or related lipid phosphate messengers. (c) 2000 Academic Press.

AN 2000:115022 HCAPLUS <<LOGINID::20091001>>

DN 132:263313

AR

ΤI Enhancement of the Migration of Metastatic Human Breast Cancer Cells by Phosphatidic Acid

ΑU Sliva, Daniel; Mason, Rebekah; Xiao, Hongyan; English, Denis

CS Experimental Cell Research Program, Methodist Research Institute, Clarian Health Partners Inc., Indianapolis, IN, 46202, USA

SO Biochemical and Biophysical Research Communications (2000). 268(2), 471-479

CODEN: BBRCA9; ISSN: 0006-291X

PB Academic Press

DТ Journal

English LA

osc.g 33 THERE ARE 33 CAPLUS RECORDS THAT CITE THIS RECORD (33 CITINGS)

RE.CNT 49 THERE ARE 49 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L23 ANSWER 16 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN

- Lysophosphatidic acid-induced Ca2+ mobilization in the neural retina of chick embryo
- Lysophosphatidic acid (LPA) plays various roles in the regulation of cell growth as a lipid mediator. We studied the effect of LPA on intracellular Ca2+ concentration ([Ca2+]i) with Fura-2 in the neural retina of chick embryo during neurogenesis. Bath application of LPA (1-100 μM) to the embryonic day 3 (E3) chick retina caused an increase in [Ca2+]i in a dose-dependent manner, with an EC50 value of 9.2 µM. The Ca2+ rise was

also evoked in a Ca2+-free medium, suggesting that release of Ca2+ from intracellular Ca2+ stores (Ca2+ mobilization) was induced by LPA. U-73122, a blocker of phospholipase C (PLC), inhibited the Ca2+ rise to LPA. Pertussis toxin partially inhibited the Ca2+ rise to LPA, indicating that Gi/Go protein was at least partially involved in the LPA response. The developmental profile of the LPA response was studied from E3 to E13. The Ca2+ rise to LPA declined drastically from E3 to E7, in parallel with decrease in mitotic activity of retinal progenitor cells. The signal transduction pathway and developmental profile of the Ca2+ response to LPA were the same as those of the Ca2+ response to ATP, which enhances the proliferation of retinal progenitor cells. The coapplication of LPA with ATP resulted in enhancement of Ca2+ rise in the E3 chick retina. Our results show that LPA induces Ca2+ mobilization in the embryonic chick retina during neurogenesis.

- AN 2000:14366 HCAPLUS << LOGINID::20091001>>
- DN 132:163687
- TI Lysophosphatidic acid-induced Ca2+ mobilization in the neural retina of chick embryo
- AU Zhou, Wen-Liang; Sugioka, Miho; Yamashita, Masayuki
- CS Department of Physiology, Osaka University Medical School, Suita, 565-0871, Japan
- SO Journal of Neurobiology (1999), 41(4), 495-504 CODEN: JNEUBZ; ISSN: 0022-3034
- PB John Wiley & Sons, Inc.
- DT Journal
- LA English
- OSC.G 6 THERE ARE 6 CAPLUS RECORDS THAT CITE THIS RECORD (6 CITINGS)
 RE.CNT 43 THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS RECORD
 ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L23 ANSWER 17 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI A metabolic path for the degradation of lysophosphatidic acid, an inhibitor of lysophosphatidylcholine
- lysophospholipase, in neuronal nuclei of cerebral cortex AB Neuronal nuclei isolated from rabbit cerebral cortex were found to be enriched in an NEM-insensitive lysophosphatidic acid (lysoPA) phosphohydrolase activity. LysoPA is an inhibitor of the nuclear lysophosphatidylcholine (lysoPC) lysophospholipase, and by preserving lysoPC levels, lysoPA boosted the nuclear production of the acyl analog of platelet-activating factor by promoting the acetylation of lysoPC (Baker, R. R.; Chang, H.-y., 1999). The nuclear phosphohydrolase converts lysoPA to 1-monoacylglycerol, and thus eliminates this lysoPA inhibition of lysoPC lysophospholipase. The nuclear lysoPA phosphohydrolase specific activity was more than three times that observed for the nuclear lysoPA lysophospholipase (Baker, R. R.; Chang, H.-y., 1999), and represents a more active route for nuclear lysoPA removal. The neuronal nuclear lysoPA phosphohydrolase was inhibited at acidic pH, and also inhibited by calcium ions. The 1-monoacylglycerol product of the phosphohydrolase is rapidly degraded by neuronal monoacylglycerol lipase, an enzyme some sevenfold more active than the phosphohydrolase and sensitive to inhibition by arachidonoyl trifluoromethyl ketone (AACOCF3). Both acidic pH and free fatty acid inhibited the lipase. In the absence of AACOCF3, production of fatty acid from lysoPA substrate could be largely attributed to the sequential actions of the nuclear phosphohydrolase and lipase. This facilitates fatty acid recycling back into phospholipid by lysophospholipid acylation when ATP levels are restored following periods of brain ischemia. At relatively low concns., sphingosine-1phosphate, and alkylglycerophosphate were the most effective phosphohydrolase inhibitors while phosphatidic acid, alkylacetylglycerophosphate and ceramide were without effect. LysoPA is an interesting regulatory mol. that can potentially preserve

lysophosphatidylcholine within the nuclear membrane for use in acetylation reactions. Thus conditions relevant to brain ischemia such as falling pH, falling ATP concns., rising fatty acid and intracellular calcium levels may, by slowing this metabolic path for lysoPA loss, promote the production of acvl PAF and contribute to the increased levels of the acetylated lipids noted in ischemia.

- AN 1999:793729 HCAPLUS <<LOGINID::20091001>>
- DN 132:149456
- TI A metabolic path for the degradation of lysophosphatidic acid, an inhibitor of lysophosphatidylcholine lysophospholipase, in neuronal nuclei of cerebral cortex
- Baker, R. R.; Chang, H.-y. ΑU
- CS Department of Biochemistry, University of Toronto, Toronto, ON, Can. SO
 - Biochimica et Biophysica Acta, Molecular and Cell Biology of Lipids (2000), 1483(1), 58-68 CODEN: BBMLFG; ISSN: 1388-1981
- Elsevier B.V. PR
- DT Journal
- LA English
- OSC.G 14 THERE ARE 14 CAPLUS RECORDS THAT CITE THIS RECORD (14 CITINGS)
- THERE ARE 48 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 48 ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L23 ANSWER 18 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- In vitro autoradiographic visualization of
- quanosine-5'-0-(3-[35s]thio)triphosphate binding stimulated by sphingosine 1-phosphate and lysophosphatidic acid
- AB Sphingosine 1-phosphate or lysophosphatidic acid activation of guanosine-5'-0-(3-[358]thio)-triphosphate ([358]GTPγ8) binding to G proteins was studied by in vitro autoradiog. in rat and guinea pig brain. The highest stimulation of [35S]GTPyS binding by sphingosine 1phosphate was observed in the mol. layer of the cerebellum. Marked stimulation was observed in most forebrain areas, including neocortex and striatum. With the exception of the substantia gelatinosa and nucleus of the solitary tract, sphingosine 1-phosphate-enhanced binding was weaker in the brainstem and spinal cord. Lysophosphatidic acid-enhanced labeling was only observed in white matter areas. The G protein inhibitor 5'-p-fluorosulfonylbenzoyl quanosine completely inhibited lysophosphatidic acid-enhanced [35S]GTPyS binding but only partially sphingosine 1-phosphate-enhanced binding. N-Ethylmaleimide abolished binding stimulated by both agonists. Sphingosine 1-phosphate enhanced labeling by another GTP analog
 - $(\beta, \gamma-imido[8-3H]guanosine-5'-triphosphate)$ similarly to that of
 - [35S]GTPyS. Lysophosphatidic acid stimulated [35S]GTPyS
 - binding in the olfactory bulb, glia limitans, and cortical subventricular zone of 1-day-old rats, whereas enhanced labeling was not observed in the latter area of 5-day-old rats. Sphingosine 1-phosphate stimulated binding in the cortical and striatal subventricular zones and
 - olfactory bulb in 1- and 5-day-old rats. In the absence of radioligand for sphingosine 1-phosphate and lysophosphatidic acid receptors, [355]GTPyS autoradiog, provides a unique opportunity to study the spatial distribution, ontogeny, and coupling properties of these
- receptors. 1999:547170 HCAPLUS <<LOGINID::20091001>> AN
- DN 131:283533
 - In vitro autoradiographic visualization of
 - quanosine-5'-0-(3-[355]thio)triphosphate binding stimulated by sphingosine 1-phosphate and lysophosphatidic acid Waeber, Christian; Chiu, Mary L.
 - AΠ
 - CS Massachusetts General Hospital and Harvard Medical School, Charlestown, MA, 02129, USA

- SO Journal of Neurochemistry (1999), 73(3), 1212-1221 CODEN: JONRA9; ISSN: 0022-3042
- PB Lippincott Williams & Wilkins
- DT Journal
- LA English
- OSC.G 16 THERE ARE 16 CAPLUS RECORDS THAT CITE THIS RECORD (16 CITINGS)
- RE.CNT 40 THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L23 ANSWER 19 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Influence of pertussis toxin on local progression and metastasis after orthotopic implantation of the human prostate cancer cell line PC-3 in nude mice
- AB Tumor cell migration is a fundamental process of metastasis. Pertussis toxin inhibits lysophosphatidic acid-related cell migration by ADP-ribosylation of G proteins. The authors examined the influence of pertussis toxin (PTX) on the progression and metastasis of the human hormone-insensitive prostate cancer cell line PC-3 after orthotopic implantation in nude mice. In 30 athymic male nude mice (NMRI), 5 + 105 PC-3 cells were injected into the dorsal prostate. After 7 d, 15 mice received a total of 6 i.p. injections of 5 µg PTX/100 g at an interval of 4 d. The other 15 mice received phosphate-buffered saline and served as control. All mice were killed at 37 d followed by macroscopical and histol, evaluation of local tumor growth and metastasis. In the control group, tumorigenicity was 100% (15 out of 15). Mean weight of the tumor-bearing unit of prostate and seminal vesicles was 541 mg (243-763 mg). PTX following orthotopic implantation of the human hormone-insensitive PC-3 cell line significantly reduces local tumor growth as well as metastasis to loco-regional lymph nodes
- AN 1999:145092 HCAPLUS <<LOGINID::20091001>>
- DN 130:333942
- TI Influence of pertussis toxin on local progression and metastasis after orthotopic implantation of the human prostate cancer cell line PC-3 in nude mice
- AU Bex, A.; Lummen, G.; Rembrink, K.; Otto, T.; Metz, K.; Rubben, H.
- CS Clinic of Urology, University of Essen Medical School, Essen, Germany
- SO Prostate Cancer and Prostatic Diseases (1999), 2(1), 36-40 CODEN: PCPDFW; ISSN: 1365-7852
- PB Stockton Press
- DT Journal
- LA English
- OSC.G 12 THERE ARE 12 CAPLUS RECORDS THAT CITE THIS RECORD (12 CITINGS)
- RE.CNT 22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L23 ANSWER 20 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
 TI Actin depolymerizing factor and cofilin phosphorylation dynamics: response
 to signals that regulate neurite extension
- AB The actin assembly-regulating activity of actin depolymg, factor (ADF)/cofilin is inhibited by phosphorylation. Studies were undertaken to characterize the signaling pathways and phosphatases involved in activating phosphorylated ADF (pADF), emphasizing signals related to neuronal process extension. Western blots using antibodies to ADF and cofilin, as well as an ADF/cofilin phosphoepitope-specific antibody characterized in this paper, were used to measure changes in the phosphorylation state and phosphate turnover of ADF/cofilin in response to inhibitors and agents known to influence growth come motility. Increases in both [Ca2+1] and cAMF levels induced rapid pADF dephosphorylation in HT4 and cortical neurons. Calcium-dependent dephosphorylation depended on the activation of protein phosphatase 2B

(PP2B), while cAMP-dependent dephosphorylation was likely through activation of PP1. Growth factors such as NGF and insulin also induced rapid pADF/pcofilin dephosphorylation, with NGF-stimulated dephosphorvlation in PC12 cells correlated with the translocation of ADF/cofilin to ruffling membranes. Of special interest was the finding that the rate of phosphate turnover on both pADF and pcofilin could be enhanced by growth factors without changing net pADF levels, demonstrating that growth factors can activate bifurcating pathways that promote both phosphorvlation and dephosphorvlation of ADF/cofilin. All exptl. results indicated that dynamics of phosphorylation on ADF and cofilin are coordinately regulated. Signals that decreased pADF levels are associated with increased process extension, while agents that increased pADF levels, such as lysophosphatidic acid, inhibit process extension. These data indicate that dephosphorylation/activation of pADF is a significant response to the activation of signal pathways that regulate actin dynamics and alter cell morphol. and neuronal outgrowth.

1998:110981 HCAPLUS <<LOGINID::20091001>> AN

DN 128:215788

OREF 128:42709a,42712a

Actin depolymerizing factor and cofilin phosphorylation dynamics: response to signals that regulate neurite extension

ΑU Meberg, Peter J.; Ono, Shoichiro; Minamide, Laurie S.; Takahashi, Masami; Bamburg, James R.

Department of Biochemistry and Molecular Biology, Colorado State CS University, Fort Collins, CO, 80523-1870, USA

SO Cell Motility and the Cytoskeleton (1998), 39(2), 172-190 CODEN: CMCYEO; ISSN: 0886-1544

Wilev-Liss, Inc.

PB DT Journal

LA English

OSC.G 127 THERE ARE 127 CAPLUS RECORDS THAT CITE THIS RECORD (127 CITINGS) RE.CNT 75 THERE ARE 75 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L23 ANSWER 21 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN

TΙ Structure/activity relationships in lysophosphatidic acid: the 2-hydroxyl moiety

AB Although lipid phosphoric acid mediators such as lysophosphatidic acid (LPA) are now recognized widely as intercellular signaling mols., the medicinal chemical of these mediators is poorly developed. With the goal of achieving a better understanding of the structure activity relationships in LPA, we have synthesized and tested a series of LPA analogs that lack the 2-hydroxyl moiety. Our series consisted of compds. with 2, 3, or 4 carbon diol or amino alc. backbones and oleovl or palmitoleovl acvl groups. These mols, cannot be acvlated further to form phosphatidic acids, nor do they have chiral centers. The rank order potency of these compds, in mobilization of calcium in MDA MB-231 cells suggested a maximum optimal chain length of 24-25 atoms. However, high potency for the inhibition of adenylyl cyclase in these cells was achieved only by one compound that also contained a dissociable proton five bond lengths from the phosphorus atom. That compound, N-oleov1-2-hydroxyethy1-1-phosphate, was nearly equipotent to 1-oleoyl LPA in both assays. The striking mimicry of LPA by the ethanolamine-based compound and the presence of fatty acid amides in tissue prompts us to propose that phosphorylated N-acyl ethanolamides occur naturally.

1997:459506 HCAPLUS <<LOGINID::20091001>> AN

DN 127:174436

OREF 127:33753a,33756a

Structure/activity relationships in lysophosphatidic acid: the 2-hydroxyl

moiety

- Lynch, Kevin R.; Hopper, Darrin W.; Carlisle, Steven J.; Catalano, John AII G.; Zhang, Ming; Macdonald, Timothy L.
- Department of Pharmacology, University of Virginia, Charlottesville, VA, 22908. USA
- Molecular Pharmacology (1997), 52(1), 75-81 SO

CODEN: MOPMA3; ISSN: 0026-895X

PB Williams & Wilkins

DT Journal LA English

OSC.G 27 THERE ARE 27 CAPLUS RECORDS THAT CITE THIS RECORD (27 CITINGS)

- L23 ANSWER 22 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- ΤI Characterization of sphingosine 1-phosphate-induced actions and its signaling pathways in rat hepatocytes
- Sphingosine 1-phosphate (S-1-P) and lysophosphatidic acid (LPA) stimulated glycogen phosphorylase, a rate-limiting enzyme responsible for glycogenolysis, in association with Ca2+ mobilization and phospholipase C (PLC) activation in rat hepatocytes. S-1-P, but not LPA, also inhibited adenosine 3',5'-cyclic monophosphate accumulation reflecting adenylyl cyclase inhibition. S-1-P-induced PLC activation, Ca2+ mobilization, and phosphorylase activation were markedly enhanced by primary culture of the cells for 24 h, whereas the inhibitory adenosine 3',5'-cyclic monophosphate response was unchanged by increasing culture time. Activation of the PLC-Ca2+ system during primary culture was specific to the lysosphingolipid; PLC and Ca2+ responses to LPA and NaF were unchanged or slightly attenuated by increasing culture time. Pertussis toxin treatment almost completely suppressed the S-1-P-induced inhibition of adenylyl cyclase but hardly influenced the lipid-induced activation of PLC and its cascade reactions. It is concluded that S-1-P, through an LPA receptor-independent mechanism, stimulates two signaling pathways, i.e., activation of the PLC-Ca2+ system and inhibition of adenylyl cyclase, through distinct S-1-P receptor-transducer systems, resulting in the modulation of glycogenolysis in rat hepatocytes.

1997:354979 HCAPLUS <<LOGINID::20091001>> AΝ

DN 127:79075

OREF 127:15101a,15104a

- Characterization of sphingosine 1-phosphate-induced actions and its signaling pathways in rat hepatocytes
- Im, Dong-Soon; Fujioka, Toshiyuki; Katada, Toshiaki; Kondo, Yoichi; Ui, AU Michio; Okajima, Fumikazu
- Lab. Signal Transduction, Inst. Mol. Cellular Regulation, Gunma Univ., Maebashi, 371, Japan

American Journal of Physiology (1997), 272(5, Pt. 1), G1091-G1099

CODEN: AJPHAP; ISSN: 0002-9513

PB DT Journal

American Physiological Society

LA English

SO

OSC.G THERE ARE 40 CAPLUS RECORDS THAT CITE THIS RECORD (40 CITINGS) 40 RE.CNT 30 THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L23 ANSWER 23 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- N-palmitoyl-serine and N-palmitoyl-tyrosine phosphoric acids are selective competitive antagonists of the lysophosphatidic acid receptors
- AB Lysophosphatidic acid is best characterized as member of a lipid mediator family with growth factor-like activities that act through a class of G protein-coupled plasma membrane receptors. In Xenopus Levis oocytes, lysophosphatidate activates at least two pharmacol. distinct receptor

subtypes distinguished by 1-acyl-sn-qlycero-2,3-cyclic phosphate Both of these naturally occurring ligands elicit oscillatory Clcurrents in the occyte through G protein-coupled activation of the phosphoinositide/Ca2+ second messenger system, which in turn leads to the opening of Ca2+ -activated C1- channels. We developed an improved chemical synthesis and purification procedure for two N-acylated amino acid phosphates. N-Palmitoyl-serine and N-palmitoyl-tyrosine phosphoric acids inhibited the lysophosphatidate-activated C1- currents with IC50 values of 5.4 and 6.5 nM at the high affinity site and 805 and 172 nM at the low affinity receptor site, resp. In selective activation of the cyclic lysophosphatidate receptor, IC50 values of 330 and 490 nM were obtained, resp. The D- and L-stereoisomers were equally effective when applied extracellularly. In contrast, they were ineffective when microinjected into the oocyte, indicating an extracellular site of inhibition. The inhibitors did not alter currents elicited by the different acetylcholine, serotonin, and glutamate receptors expressed heterologously in the oocyte. Pharmacol. anal. of the results indicates that N-palmitoyl-serine and N-palmitoyl-tyrosine phosphoric acids are potent and specific competitive inhibitors of the lysophosphatidate receptors in the X. Levis oocyte. 1996:570299 HCAPLUS <<LOGINID::20091001>> 125:265929 OREF 125:49377a,49380a N-palmitovl-serine and N-palmitovl-tyrosine phosphoric acids are selective competitive antagonists of the lysophosphatidic acid receptors Liliom, Karoly; Bittman, Robert; Swords, Bernadette; Tigyi, Gabor Department of Physiology and Biophysics, University of Tennessee, Memphis, TN, 38163, USA Molecular Pharmacology (1996), 50(3), 616-623

CODEN: MOPMA3; ISSN: 0026-895X PB Williams & Wilkins

DT Journal LA English

AN DN

AII CS

SO

OSC.G 56 THERE ARE 56 CAPLUS RECORDS THAT CITE THIS RECORD (56 CITINGS)

L23 ANSWER 24 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN Inhibitors of lipid phosphatidate receptors: N-palmitoyl-serine and

N-palmitoyl-tyrosine phosphoric acids

AB An improved synthesis of two lipid phosphoric acids, N-palmitovl-L-serinephosphoric acid (NP-Ser-PA) and N-palmitovl-L-tvrosinephosphoric acid (NP-Tyr-PA), from L-serine and L-tyrosine benzyl esters is described. The sequence of N-acylation, followed by phosphitylation with dibenzyl N, N-diisopropylphosphoramidite, oxidation to the corresponding phosphate triesters, and simultaneous debenzylation of the dibenzyl phosphate and benzyl carboxylic esters gave NP-Ser-PA and NP-Tyr-PA in high overall yields. NP-Ser-PA and NP-Tyr-PA and their D-stereoisomers were potent reversible inhibitors of the lysophosphatidic acid receptors expressed in Xenopus oocytes, thus providing prototypic structures for the development of inhibitors of the lysophosphitidate family of phospholipid growth factors.

1996:154614 HCAPLUS <<LOGINID::20091001>> AN

DN 124:317814

OREF 124:58961a,58964a

Inhibitors of lipid phosphatidate receptors: N-palmitoyl-serine and N-palmitoyl-tyrosine phosphoric acids

AII Bittman, Robert; Swords, Bernadette; Liliom, Karoly; Tigyi, Gabor

CS Dep. Chemistry Biochemistry, Queens College City Univ. New York, Flushing, NY, 11367-1597, USA

SO Journal of Lipid Research (1996), 37(2), 391-8 CODEN: JLPRAW; ISSN: 0022-2275

- PB Lipid Research, Inc.
- DT Journal LA English
- OSC.G 24 THERE ARE 24 CAPLUS RECORDS THAT CITE THIS RECORD (24 CITINGS)
- L23 ANSWER 25 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Selective inhibition of DNA polymerase- α family with chemically
- synthesized derivatives of PHYLPA, a unique Physarum lysophosphatidic acid AB PHYLPA, a unique Physarum lysophosphatidic acid (LPA),
 - showed selective inhibition of a family of DNA polymerase α , including DNA polymerases α , δ and ϵ ; but no inhibition of DNA polymerase β or γ was observed. To reveal the mol. mechanism of inhibition of DNA polymerases by PHYLPA, four stereoisomers and some other derivs. were synthesized and their effects on DNA polymerases were studied. Among eight derivs. synthesized, PHYLPA-1 (the natural PHYLPA; sodium 1-O-[(9'S,10'R)-9',10'-methanohexadecanoy1]-snglycerol 2,3-cyclic phosphate) and PHYLPA-2 (sodium 3-0-[(9'S,10'R)-9',10'-methanohexadecanoy1]-sn-glycerol 1,2-cyclic phosphate) were strong and specific inhibitors of a family of DNA polymerase α. But their stereoisomers PHYLPA-3 (sodium 1-0-((9'R,10'S)-9',10'-methanohexadecanovl)-sn-glycerol 2,3-cyclic phosphate) and PHYLPA-4 (sodium 3-0-[(9'R,10'S)-9',10'-methanohexadecanovl]-sn-glycerol 1,2-cyclic phosphate) were weak inhibitors, showing the critical importance of stereochem. of a cyclopropane-containing fatty acid for the inhibitory activity. Some derivs. having no cyclopropane-containing fatty acids palmitoyl-, oleoyl-, and palmitoleoyl-PHYLPA -showed inhibition to some extent; but 1-palmitoyl and 1-oleoyl lysophosphatidic acid, which has no cyclic phosphate, did not show an apparent inhibitor activity on DNA polymerases. Hence, the extent of the inhibition apparently depends on the stereochem. of both the fatty acid moiety and the cyclic
- phosphate.
 AN 1995:764871 HCAPLUS <<LOGINID::20091001>>
- DN 123:221508
- OREF 123:39331a,39334a
- TI Selective inhibition of DNA polymerase- α family with chemically
- synthesized derivatives of PHYLPA, a unique Physarum lysophosphatidic acid AU Murakami-Murofushi, Kimikok, Kobayashi, Susumu, Onimura, Kenjiro, Matsumoto, Miyoko; Shioda, Masaki; Yoshida, Shonen; Shoji, Mami; Murofushi, Hirom
- CS Department of Biology, Faculty of Science, Ochanomizu University, Ohtsuka 2-1-1, Bunkyo-ku, Tokyo, 112, Japan
- SO Biochimica et Biophysica Acta, Lipids and Lipid Metabolism (1995), 1258(1), 57-60 CODEN: BBLLA6; ISSN: 0925-4439
- PB Elsevier B.V. DT Journal
- DI JOULNAL
- LA English
- OSC.G 8 THERE ARE 8 CAPLUS RECORDS THAT CITE THIS RECORD (8 CITINGS)
- L23 ANSWER 26 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- ${\tt TI} \quad {\tt Use \ of \ photoreactive \ substrates \ for \ characterization \ of \ lysophosphatidate} \\ acyltransferases \ from \ developing \ soybean \ cotyledons$
- AB Photoreactive lipid analogs, namely,
 - 1-acyl-2-(12-azidooleoyl)glycero-3-phosphocholine (N3-PC) and 1-acyl-2-(12-azidooleoyl)glycero-3-phosphoethanolamine (N3-PE) have been synthesized as previously described [R. Rajasekharan and J. D. Kemp (1994) J. Lipid Res. 35, 45-51]. Azidophosphatidic acid was produced by hydrolyzing N3-PC with phospholipase D. All of the lysophospholipid analogs, 2-(12-azidooleoyl)glycero-3-phosphate (N3-LPA),

2-(12-azidooleoyl)glycero-3-phosphocholine (N3-LPC), and 2-(12-azidooleoyl)glycero-3-phosphoethanolamine (N3-LPE), were produced from appropriate azidophospholipids by lipase treatment. The photoactive lysophospholipid analogs were recognized as substrates by acyltransferases in the dark and as irreversible inhibitors after photolysis with UV light. The photoinactivation of acyltransferases by azidolysophospholipids was protected by the addition of natural lysophospholipids. Incubation of developing soybean microsomal membranes with N3-LPA followed by photolysis resulted in 69% inhibition of lysophosphatidic acid (LPA) acyltransferase and also had significant inhibitory effects on lysophosphatidylcholine (LPC) and lysophosphatidylethanolamine (LPE) acyltransferases, indicating that the LPA analog interacts with all

acid (LPA) acyltransferase and also had significant inhibitory effects on lysophosphatidylcholine (LPC) and lysophosphatidylcholine (LPC) and lysophosphatidylcholine (LPC) and lysophosphatidylethanolamine (LPE) acyltransferases, indicating that the LPA analog interacts with all the lysophospholipid acyltransferases. When the membranes were photolyzed with N3-LPC or N3-LPE and assayed, the membranes showed approx. 50% inactivation of LPC and LPE acyltransferase activities, whereas LPA acyltransferase was unaffected, suggesting that a single enzyme might acyltate both LPC and LPE. The recognition of these photoreactive lipid analogs by acyltransferases will facilitate the identification and purification of these membrane-bound enzymes.

- AN 1994:599203 HCAPLUS <<LOGINID::20091001>>
- DN 121:199203
- OREF 121:36094h,36095a
- TI Use of photoreactive substrates for characterization of lysophosphatidate acyltransferases from developing soybean cotyledons
- AU Rajasekharan, Ram; Nachiappan, Vasanthi
- CS Plant Genetic Eng. Lab., New Mexico State Univ., Las Cruces, NM, 88003, USA
- SO Archives of Biochemistry and Biophysics (1994), 311(2), 389-94 CODEN: ABBIA4; ISSN: 0003-9861
- DT Journal
- LA English
- OSC.G 13 THERE ARE 13 CAPLUS RECORDS THAT CITE THIS RECORD (13 CITINGS)
- L23 ANSWER 27 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- II Phosphatidic acid, lysophosphatidic acid, and lipid A are inhibitors of glycosylphosphatidylinositol-specific phospholipase D. Specific inhibition of a phospholipase by product analogs?
- AB Previous work has suggested that the glycosylphosphatidylinositol-specific phospholipase D (GPI-PLD) purified from bovine serum is inhibited by phosphatdic acid (PA). In this study, the specificity and mechanism of this phenomenon using [3H]mvristate-labeled variant surface glycoprotein dispersed in Nonidet P-40 as substrate is reported. Inhibition of GPI-PLD by PAs (IC50 .apprx.1 µM) was relatively independent of the length or degree of unsatn. of the fatty acyl chains. It was also observed that lysophosphatidic acid and several natural and synthetic lipid A prepns. were inhibitory in the same concentration range. The inhibitory potency of PA, lysophosphatidic acid, and lipid A was dependent on the detergent concentration in the assay but in all cases this was in a large (i.e. > 100-fold) molar excess over the inhibitor. The inhibitory lipids did not affect substrate availability nor did they reduce hydrolysis of variant surface glycoprotein by a bacterial phosphatidylinositol-specific phoshpolipase C. Studies with a wide range of other lipids, detergents, and phosphate esters indicated that inhibition was specific for lipids containing a phosphomonoester group. The data suggest that inhibition is due to a direct interaction between PA (or lipid A) and the GPI-PLD rather than an indirect effect on the substrate particle.
- AN 1993:250409 HCAPLUS <<LOGINID::20091001>>
- DN 118:250409

- TI Phosphatidic acid, lysophosphatidic acid, and lipid A are inhibitors of glycosylphosphatidylinositol-specific phospholipase D. Specific inhibition of a phospholipase by product analogs?
- AU Low, Martin G.; Huang, Kuo Sen
- CS Coll. Physicians Surg., Columbia Univ., New York, NY, 10032, USA
- SO Journal of Biological Chemistry (1993), 268(12), 8480-90 CODEN: JBCHA3; ISSN: 0021-9258
- OT Journal
- LA English
- OSC.G 17 THERE ARE 17 CAPLUS RECORDS THAT CITE THIS RECORD (17 CITINGS)
- L23 ANSWER 28 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Inhibition of eukaryotic DNA polymerase α with a novel lysophosphatidic acid (PHYLPA) isolated from myxoamebae of Physarum polycephalum
- AB À specific inhibitor of DNA polymerase-α was isolated from the lipid fraction prepared from mysoameobae of a true slime mold, P. polycephalum. The purified substance was subjected to structural studies by fast atom bombardment mass spectroscopy, IR spectroscopy, and 2-dimensional NMR spectroscopy. The structure of this substance was thereby suggested to be a novel lysophosphatidic acid (IPA) composed of cyclic phosphate and cyclopropane-containing hexadecanoic acid. This substance was named PHYLPA (Physarum LPA). PHYLPA inhibited >80% of affinity-purified calf thymus I activity at a concentration of 10 mg/mL (apprx.20 μM). Inhibition was observed for I but not for DNA polymerase I from various eukaryotic species, nor did it inhibit DNA polymerase I from E. coli. From kinetic analyses, the inhibition was considered to be caused by the interaction of PHYLPA with template DNA.
- AN 1992:607756 HCAPLUS <<LOGINID::20091001>>
- DN 117:207756
- OREF 117:35761a,35764a
- TI Inhibition of eukaryotic DNA polymerase α with a novel lysophosphatidic acid (PHYLPA) isolated from myxoamebae of Physarum polycephalum
- AU Murakami-Murofushi, Kimiko; Shioda, Masaki; Kaji, Kazuhiko; Yoshida, Shonen; Murofushi, Hiromu
- CS Fac. Sci., Ochanomizu Univ., Tokyo, 112, Japan
- SO Journal of Biological Chemistry (1992), 267(30), 21512-17
- CODEN: JBCHA3; ISSN: 0021-9258
- DT Journal LA English
- OSC.G 33 THERE ARE 33 CAPLUS RECORDS THAT CITE THIS RECORD (34 CITINGS)
- L23 ANSWER 29 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Characterization of dolichol and dolichyl phosphate phosphatase from sova beans (Glycine max)
- AB A series of polyprenols, ranging in length from 15 to 22 isoprene units, has been isolated from soybeans (Glycine max) and purified by high-pressure liquid chromatog. NMR, IR, and mass spectra of the compds. indicated that they are α-saturated polyprenols of the dolichol type. The amount present in dry seeds was .appxx. 9 mg/100 g, whereas dolichyl phosphate (Dol-P) was present only in trace amts. Dol-P phosphatase (DPP) activity was detected in the microsomal fraction of 5-day-old germinating soybean cotyledons. The DPP activity was linear with respect to time and protein concentration and exhibited a broad pH optimum (pH 7-9). Triton X-100 was necessary for significant enzyme activity. Enzyme activity was slightly enhanced by EDTA, whereas dithiothreitol was without effect. An apparent Km of 5 μM was determined for Dol-P. Bivalent metal ions were not required for enzyme activity. A number of phosphorylated compds, tested as enzyme substrates (including a number of nucleoside

phosphates, glucose 6-phosphate, Na β-glycerophosphate, and Na4P2O7) did not compete with [1-3H]Do1-P as substrate. A number of phospholipids were also tested for their ability to act as DPP substrates. At 1 mM concentration, phosphatidylcholine, phosphatidylethanolamine, phosphatidic acid, and lysophosphatidic acid each exhibited enzymic activity. However, at 0.1 mM concentration, phosphatidylcholine and phosphatidylethanolamine were slightly stimulatory, whereas phosphatidic acid and lysophosphatidic acid were still inhibitory. Phosphatidic acid showed competitive inhibition.

AN 1983:485171 HCAPLUS <<LOGINID::20091001>>

DN 99:85171

OREF 99:13097a,13100a

- TΙ Characterization of dolichol and dolichyl phosphate phosphatase from sova beans (Glycine max)
- ΑΠ Ravi, Kothapalli; Rip, Jack W.; Carroll, Kenneth K.
- CS Dep. Biochem., Univ. West. Ontario, London, ON, N6A 5C1, Can. Biochemical Journal (1983), 213(2), 513-18 so
 - CODEN: BIJOAK; ISSN: 0306-3275
- Journal

LA English

- OSC.G THERE ARE 6 CAPLUS RECORDS THAT CITE THIS RECORD (6 CITINGS) 6
- L23 ANSWER 30 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- ΤI Pulmonary phosphatidic acid phosphatase. A comparative study of the aqueously dispersed phosphatidate-dependent and membrane-bound phosphatidate-dependent phosphatidic acid phosphatase activities of rat
- AB The properties of the aqueously dispersed phosphatidate-dependent phosphatidate phosphatase (EC 3.1.3.4) (I) activities of rat lung were studied in microsomal and cytosol prepns. and compared with those of the membrane-bound phosphatidate-dependent activities. Microsomal I displayed a prominent pH optimum at 6.5 with a minor peak which varied between 7.5-8 in different expts. The major cytosol I activity was at the higher pH (7.5-8.0) but a distinct optimum was also observed at pH 6.0-6.5. With the membrane-bound substrate, a single broad optimum was observed between pH 7.4 and 8.0 with the cytosol and 6.5-7.5 with the microsomal fraction. Subcellular fractionation studies revealed that the microsomal fraction possessed the greatest proportion of the total I activity and the highest relative specific activity. However, studies with marker enzymes indicated that the aqueously dispersed phosphatidate-dependent activity could be present in plasma membrane, lysosomes, and osmiophilic lamellar bodies as well as in the endoplasmic reticulum. The aqueously dispersed phosphatidic acid-dependent activities present in the microsomal and supernatant fractions were inhibited by Ca2+, Mn2+, F-, and by high concns, of Mg2+. In contrast to the membrane-bound phosphatidate-dependent activities, there was little Mg2+ stimulation and only a very slight inhibitory effect was noted with EDTA. A small EDTA-dependent Mg2+ stimulation could be observed with the microsomal fraction but only at the lower pH optimum (6.5). The presence of a number of phosphate esters tended to stimulate rather than inhibit the microsomal activity, indicating that I is relatively specific for lipid substrates. Marked inhibitions were noted with lysophosphatidic acid and phosphatidylglycerol phosphate. Phosphatidylcholine produced a slight inhibition. The results indicate that the bulk of the aqueously dispersed phosphatidate-dependent I activities of rat lung microsomes and cytosol is not related to the activities observed with membrane-bound phosphatidate. The Mg2+-dependent I activities may be synonymous. However, unequivocal conclusions will only be possible when the polypeptide or polypeptides responsible for these activities can be purified.

- DN 91:153340
- OREF 91:24677a,24680a
- TI Pulmonary phosphatidic acid phosphatase. A comparative study of the aqueously dispersed phosphatidate-dependent and membrane-bound phosphatidate-dependent phosphatidic acid phosphatase activities of rat lung
- AU Yeung, Alex; Casola, Paul G.; Wong, Ching; Fellows, J. Fraser; Possmayer, Fred
- CS Dep. Obstet. Gynaecol., Univ. Western Ontario, London, ON, N6A 5A5, Can.
- SO Biochimica et Biophysica Acta, Lipids and Lipid Metabolism (1979), 574(2), 226-39
 - CODEN: BBLLA6; ISSN: 0005-2760 Journal
- DT Journa
- LA English
- OSC.G 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)
- L23 ANSWER 31 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Involvement of guanosine 5'-diphosphate-3'-diphosphate in the regulation of phospholipid biosynthesis in Escherichia coli. Lack of ppGpp inhibition of acyl transfer from acyl-ACP to sn-glycerol 3phosphate
- AB The response of the E. coli sn-qlycerol-3-phosphate acyltransferase to ppGpp has been determined in vitro employing palmityl-CoA and palmityl-ACP as acyl substrates. Levels of ppGpp which cause significant inhibition of enzyme activity with palmityl-CoA as substrate had no effect on enzyme activity when palmityl-ACP was employed as acyl donor. The inhibition of enzyme activity observed with palmityl-CoA as acyl substrate was dependent upon the relative concens of MgCl2 and ppGpp employed. With palmityl-CoA as acyl donor, ppGpp inhibited the production of lysophosphatidic acid but not phosphatidic acid. With palmityl-ACP as acyl substrate, ppGpp had no influence upon the distribution of the reaction products.
- AN 1975:493602 HCAPLUS <<LOGINID::20091001>>
- DN 83:93602
- OREF 83:14693a,14696a
- TI Involvement of guanosine 5'-diphosphate-3'-diphosphate in the regulation of phospholipid biosynthesis in Escherichia coli. Lack of ppGpp inhibition of acyl transfer from acyl-ACP to sn-glycerol 3phosphate
- AU Lueking, Donald R.; Goldfine, Howard
- CS Sch. Med., Univ. Pennsylvania, Philadelphia, PA, USA
- SO Journal of Biological Chemistry (1975), 250(13), 4911-17 CODEN: JBCHA3; ISSN: 0021-9258
- DT Journal
- LA English
- OSC.G 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD (3 CITINGS)
- L23 ANSWER 32 OF 32 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Partial purification and properties of an acyl coenzyme A:sn-glycerol 3phosphate acyltransferase from rat liver mitochondria
- AB The partial purification (6-fold) and properties of a position and substrate specific acyl coenzyme A:sn-qlycerol-3-phosphate acyltransferase from rat liver mitochondria are described. The preparation was devoid of acyl-CoA:monoacylglycerol-3-phosphate acyltransferase and lipid phosphomonoesterase activity. All of the glycerol-3-phosphate acylated in the presence of palmityl-CoA was identified as 1-palmityl-sn-glycerol-3-phosphate. The order of effectiveness of various acyl-CoA donors was palmityl > stearyl .simeq. myristyl > decanyl-CoA. Oleyl- and linoleyl CoA were .appxx. 5% as effective as palmityl CoA. Palmitic acid was esterified exclusively in position l of the sn-glycerol mol. The activity was stimulated by phosphatidylserine,

asolectin, and lecithin, whereas cardiolipin, lysophosphatidic acid, and phosphatidic acid were inhibitory. Mg2+, Ca2+, Mn2+, and to a lesser extent Co2+ enhanced the activity. The findings demonstrated that the acylation of sn-glycerol-3phosphate involves an enzyme activity sep. from that which acylates 1-palmityl-sn-qlycerol-3-phosphate. The enzyme preparation offers a convenient and efficient method for the preparation of 1-palmityl-sn-glycerol-3-phosphate.

1973:401843 HCAPLUS <<LOGINID::20091001>> AN

DN 79:1843

OREF 79:347a,350a

TΙ Partial purification and properties of an acyl coenzyme A:sn-glycerol 3phosphate acyltransferase from rat liver mitochondria

Monroy, Gladys; Kelker, Hanna Chroboczek; Pullman, Maynard E. AII

CS Public Health Res. Inst., City of New York, Inc., New York, NY, USA SO.

Journal of Biological Chemistry (1973), 248(8), 2845-52 CODEN: JBCHA3: ISSN: 0021-9258

DT Journal

English LA

OSC.G 14 THERE ARE 14 CAPLUS RECORDS THAT CITE THIS RECORD (14 CITINGS)

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L24 1 L7

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- L24 ANSWER 1 OF 1 HCAPLUS COPYRIGHT 2009 ACS on STN
- Catalysis of gas hydrates by biosurfactants in seawater-saturated sand/clav
- Biosurfactants catalyzed natural gas hydrate formation in sand/clay packs saturated with seawater. Representative samples from the five possible biosurfactant classifications enhanced hydrate formation rate and decreased hydrate induction time. Biosurfactants increased rates 96% to 288% and decreased induction times 20% to 71% relative to the control. Micellar-forming rhamnolipid reached a critical micellar concentration at 13

ppm at

hydrate-forming conditions; these micelles migrated readily through a seawater-saturated sand pack to catalyze hydrate formation in another zone. The type of biosurfactant, in conjunction with specific porous media, help determine massive, dispersed, nodular, or stratified forms of hydrates. Results suggested that minimal microbial activity in ocean-floor sands can

greatly influence gas hydrate formation.

2004:82258 HCAPLUS <<LOGINID::20091001>>

DN 140:377525

AN

TT Catalysis of gas hydrates by biosurfactants in seawater-saturated sand/clav

Rogers, Rudy E.; Kothapalli, Chandra; Lee, May S.; Woolsey, J. Robert AU

Swalm School of Chemical Engineering, Mississippi State University, MS,

SO Canadian Journal of Chemical Engineering (2003), 81(5), 973-980

CODEN: CJCEA7; ISSN: 0008-4034

PR Canadian Society for Chemical Engineering

DT Journal LA English

тт 685090-09-9

RL: CAT (Catalyst use); USES (Uses)

(catalysis of natural gas hydrates formation by biosurfactants in seawater-saturated sand/clay)

RM 685090-09-9 HCAPLUS

CN Hexadecanoic acid, (1S)-1-[[[[[[(1S)-1-carboxy-2hydroxyethyl]amino]oxy]hydroxyphosphinyl]oxy]methyl]-1,2-ethanediyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

osc.g THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD (2 CITINGS) RE.CNT 28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

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		IN FRIEDLIN)/CN
E2	2	SERINE PHENYLTHIOHYDANTOIN/CN
E3		SERINE PHOS/CN
E4	2	SERINE PHOSPHATASE/CN
E5	1	SERINE PHOSPHATASE (BACILLUS LICHENIFORMIS STRAIN ATCC 14580 GENE RSBU)/CN
E6	1	SERINE PHOSPHATASE (BACILLUS LICHENIFORMIS STRAIN ATCC 14580 GENE RSBX)/CN
E7	1	SERINE PHOSPHATASE (BACILLUS LICHENIFORMIS STRAIN ATCC 14580
		GENE SPOILE)/CN
E8	1	SERINE PHOSPHATASE (BACILLUS SUBTILIS GENE SPOIIE)/CN
E9	1	SERINE PHOSPHATASE (DEPHOSPHORYLATION OF RSBS) (BACILLUS SUB
		TILIS GENE RSBX)/CN
E10	1	SERINE PHOSPHATASE (DEPHOSPHORYLATION OF RSBV) (BACILLUS SUB- TILIS GENE RSBU)/CN
E11	5	SERINE PHOSPHATASE (FRANKIA STRAIN CCI3)/CN
E12	ī	SERINE PHOSPHATASE (GEOBACILLUS THERMODENITRIFICANS STRAIN N
	-	G80-2)/CN
=> exp serin	e phos	phate/cn
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=> s e5		
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=> exp serin	e phos	phoric/cn
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E3		SERINE PHOSPHORIC/CN
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E6	1	SERINE PROTEASE (ACINETOBACTER STRAIN ADP1)/CN

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E8	1 SERINE PROTEASE (ACREMONIUM STRAIN F11177 ISOFORM AS-	E2 FRAG
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E10	1 SERINE PROTEASE (AEROMONAS HYDROPHILA HYDROPHILA STRA 7966)/CN	IN ATCC
E11	1 SERINE PROTEASE (AEROMONAS SALMONICIDA SUBSP. SALMONI NE ASPA)/CN	CIDA GE
E12	1 SERINE PROTEASE (AGROBACTERIUM TUMEFACIENS STRAIN C58 TU4566)/CN	GENE A

=> file hcaplus COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION FILL ESTIMATED COST 5.83 6.05

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FILE COVERS 1907 - 1 Oct 2009 VOL 151 ISS 14 FILE LAST UPDATED: 30 Sep 2009 (20090930/ED) REVISED CLASS FIELDS (/NCL) LAST RELOADED: Aug 2009 USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Aug 2009

HCAplus now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2009.

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This file contains CAS Registry Numbers for easy and accurate substance identification.

The ALL, BIB, MAX, and STD display formats in the CA/Caplus family of databases have been updated to include new citing references information. This enhancement may impact record import into database management software. For additional information, refer to NEWS 9.

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       1171903 THU/RL
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                 (L1 (L) THU/RL)
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 - 09032 ATHEROSCIEROSI.
 - 8045 STENT
- 131238 CARDIOVASCULAR
- L3 194608 NEOINTIM? OR ATHEROSCLEROSIS OR STENT OR CARDIOVASCULAR
- => s 12 and 13
- L4 0 L2 AND L3
- => s 12 and (PY<2003 or AY<2003 or PRY<2003)
- 22985329 PY<2003
 - 4511505 AY<2003
 - 3981271 PRY<2003
- L5 31 L2 AND (PY<2003 OR AY<2003 OR PRY<2003)
- => d 15 1-31 ti
- L5 ANSWER 1 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Antibody-label complexes and methods for antigen or ligand immunolabeling or detection, diagnosis and therapy
- L5 ANSWER 2 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Method for purification of naturally phosphorylated peptide micelle and its uses
 - L5 ANSWER 3 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Human cDNAs encoding separase, methods for modulation of separase activity in sister chromatid DNA separation, and uses thereof
- L5 ANSWER 4 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Human G protein-coupled receptor kinase gene 69087, nuclear protein gene 15821, and protein kinase phosphatase gene 15418 and their uses
- L5 ANSWER 5 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Vaccines comprising hydrophobic liquid carrier, liposome, antigen and adjuvant
- L5 ANSWER 6 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Composition and method for the repair and regeneration of cartilage and other tissues based on a polymer gel
- L5 ANSWER 7 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI The antianxiety-like effects of antagonists of group I and agonists of group II and III metabotropic glutamate receptors after intrahippocampal administration
 - 5 ANSWER 8 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- ${\tt TI} \quad {\tt Methods} \ {\tt for} \ {\tt the} \ {\tt detection} \ {\tt of} \ {\tt modified} \ {\tt peptides}, \ {\tt proteins} \ {\tt and} \ {\tt other} \ {\tt molecules}$
- L5 ANSWER 9 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Quantitative amino acid analysis using a Beckman system gold HPLC 126AA analyzer
- L5 ANSWER 10 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- II Elevated levels of group-III metabotropic glutamate receptors in the inferior colliculus of genetically epilepsy-prone rats following intracollicular administration of L-serine-O-phosphate
- L5 ANSWER 11 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN

- In situ crosslinking of proteins for wound sealant
- 1.5 ANSWER 12 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- Compounds for inhibiting diseases and preparing cells for transplantation
- L5 ANSWER 13 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- Compositions and methods for treating amyloidosis TI
- ANSWER 14 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN L5
- TΙ Phosphocholine surfactants and their use
- ANSWER 15 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TΙ Associates of macromolecules and complex aggregates for improved payload and controlled drug delivery
- ANSWER 16 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- ΤI Methods and compositions to treat glycosaminoglycan-associated molecular interactions
- ANSWER 17 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- ΤI Biocompatible composite material
- ANSWER 18 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- ΤI Sphingolipid derivatives, their preparation, and their therapeutic use
- ANSWER 19 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- ТT Liquid compositions for disinfection of contact lenses based on Polyquaternium compounds
- ANSWER 20 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN L5
- TΙ Biomimetic calcium phosphate implant coatings and methods for making the same
- ANSWER 21 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN L5
- TI Oral drug delivery compositions comprising modified amino acids and bioactive peptides
- L5 ANSWER 22 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN Inhibiting undesirable taste in oral compositions
- ΤI
- ANSWER 23 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN 1.5
- TΙ Synthetic phosphopeptides for treating bone diseases
- L5 ANSWER 24 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TΙ Prolonged anticonvulsant action of glutamate metabotropic receptor agonists in inferior colliculus of genetically epilepsy-prone rats
- ANSWER 25 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN L5
- Effect of sterical stabilization on macrophage uptake in vitro and on ΤI thickness of the fixed aqueous layer of liposomes made from alkylphosphocholines
- ANSWER 26 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN L5
- Activation of group III metabotropic glutamate receptors is neuroprotective in cortical cultures
- L.5 ANSWER 27 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- ΤТ Diketopiperazine-based drug delivery systems
- 1.5 ANSWER 28 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Antitumor liposomes containing phospholipid analogs and ether lipids

- L5 ANSWER 29 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Mycobacterium-derived organic phosphate compounds as activators of Ty δ lymphocytes
- L5 ANSWER 30 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Modification of implant surface with bioactive conjugates for improved integration into tissue
- L5 ANSWER 31 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Drug preparations of reduced toxicity

=> file stnguide

COST IN U.S. DOLLARS SINCE FILE TOTAL SENTRY SESSION 14.63 20.68

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FILE CONTAINS CURRENT INFORMATION. LAST RELOADED: Sep 25, 2009 (20090925/UP).

=> d 15 5 6 10 11 12 13 14 17 18 23 ti abs bib YOU HAVE REQUESTED DATA FROM FILE 'HCAPLUS' - CONTINUE? (Y)/N:y

- L5 ANSWER 5 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Vaccines comprising hydrophobic liquid carrier, liposome, antigen and adjuvant
- AB The present invention is concerned with vaccines and their preparation An effective long-term immune response, especially in mammals, can be produced using a vaccine comprising an antigen encapsulated in liposomes, a suitable adjuvant and a carrier comprising a continuous phase of a hydrophobic substance. The vaccine is particularly effective in eliciting the production of antibodies that recognize epitopes of native proteins. The antigen is viral, bacterial, protozoal or mammalian antigen such as zona pellucida, alc. dehydrogenase, hepatitis B or streptokinase; the liposome comprises unesterified cholesterol and a phospholipid selected from phosphogiosteril, phosphochanolamine, phosphoserine, phosphocholine and phosphoinositol; the hydrophobic liquid carrier is an oil (mineral oil, vegetable oil or nut oil) or water-in-oil emulsion; and the adjuvant is alum or aluminum compound or TiterMax. A long-term immunocontraceptive for mammal comprising zona pellucida is disclosed.
- AN 2002:368338 HCAPLUS <<LOGINID::20091001>>
- DN 136:368452
- TI Vaccines comprising hydrophobic liquid carrier, liposome, antigen and adjuvant
- IN Brown, Robert George; Pohajdak, William; Kimmins, Warwick Charles
- PA Immunovaccine Technologies Inc., Can.
- SO PCT Int. Appl., 66 pp. CODEN: PIXXD2
- DT Patent
- LA English
- FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OSC.G 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD (3 CITINGS)
RE.CNT 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L5 ANSWER 6 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
- TI Composition and method for the repair and regeneration of cartilage and other tissues based on a polymer gel
- AB The present invention relates to a new method for repairing human or animal tissues such as cartilage, meniscus, ligament, tendon, bone, skin, cornea, periodontal tissues, abscesses, resected tumors, and ulcers. The method comprises the step of introducing into the tissue a temperature-dependent

polymer gel composition such that the composition adhere to the tissue and promote

support for cell proliferation for repairing the tissue. Other than a polymer, the composition preferably comprises a blood component such as whole blood, processed blood, venous blood, arterial blood, blood from bone, blood from bone—marrow, bone marrow, umbilical cord blood, placenta blood, erythrocytes, leukocytes, monocytes, platelets, fibrinogen, thrombin and platelet rich plasma. The present invention also relates to a new composition to be used with the method of the present invention. For example, chondral defects with perforations to the subchrondal bone of rabbits were treated with a peripheral blood/chitosan-qlyceryl phosphate mixture that was delivered as a liquid, and allowed to solidify in situ. After 5-8 wk healing, the blood/chitosan-treated defects were filled with repair tissue having the appearance of hyaline, a glycosaminoglycan (GAG)-rich cartilage repair tissue, which adhered to the defects surfaces, and filled the defects. Repair tissue from the untreated defects (control) had the

appearance of fibro-cartilage, with particularly no metachromatic staining for GAG, and only partial defect filling.

AN 2002:10323 HCAPLUS <<LOGINID::20091001>>

- 136:74708 DN
- TI Composition and method for the repair and regeneration of cartilage and other tissues based on a polymer gel
- IN Hoemann, Caroline D.; Buschmann, Michael D.; Mckee, Marc D.
- PA Biosyntech Canada Inc., Can.
- SO PCT Int. Appl., 106 pp. CODEN: PIXXD2
- DT Patent
- LA English
- FAN CHT 1

FAN.			NO.		KIND DATE						APPL	ICAT	ION	DATE						
PI	WO WO	2002	0002 0002	72 72		A2 20020103 A3 20020808					WO 2	001-	CA95							
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	US	2001 2005	-313	25		A1		2005	0107											

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT ALL CITATIONS AVAILABLE IN THE RE FORMAT

OSC.G 15 RE.CNT 7 THERE ARE 15 CAPLUS RECORDS THAT CITE THIS RECORD (20 CITINGS) THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD

1.5 ANSWER 10 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN

TI Elevated levels of group-III metabotropic glutamate receptors in the inferior colliculus of genetically epilepsy-prone rats following

intracollicular administration of L-serine-O-phosphate AB The selective group-III metabotropic glutamate receptor agonist, L-serine-O-phosphate (L-SOP), when injected bilaterally into the inferior colliculus of the sound sensitive genetically epilepsy-prone (GEP) rats produces a short proconvulsant excitation followed by a long phase of protection against sound-induced seizures lasting for 2-4 days. We have studied this prolonged suppression of audiogenic seizures using pharmacol. and mol. biol. approaches including semiguant. RT-PCR and western blotting. The intracerebroventricular injection of the protein synthesis inhibitor cycloheximide (120 µg) 30 min beforehand significantly reduces the proconvulsant seizure activity and the prolonged anticonvulsant effect of intracollicular L-SOP (500 nmol/side). The sensitive semiquant. RT-PCR revealed a significant up-regulation in mGlu4 and mGlu7 mRNA levels in the inferior colliculus at 2 days (maximum suppression of audiogenic seizures) after intracollicular L-SOP injection compared with the non-injected, 2-day post-vehicle treated and 7-day (return to expressing audiogenic seizures) post-drug or vehicle-treated groups. No significant changes were observed in mGlu6 or mGlu8 mRNA expression levels in drug-treated compared with control groups. Examination of mGlu4a and mGlu7a protein levels using western blotting showed a significant increase in mGlu7a but no significant change in mGlu4a protein levels 2 days after L-SOP treatment compared with the control groups (non-injected and 2-day vehicle-injected group). These results suggest that up-regulation of mGlu7 receptors is involved in the prolonged anticonvulsant effect of L-SOP against sound-induced seizures in GEP rats.

The potential use of mGlu7 agonists as novel anti-epileptic agents merits

AN 2001:508586 HCAPLUS <<LOGINID::20091001>>

DN 135:298653

investigation.

TI Elevated levels of group-III metabotropic glutamate receptors in the inferior collicular of genetically epilepsy-prone rats following intracollicular administration of L-serine-O-phosphate

AU Yip, Ping K.; Meldrum, Brian S.; Rattray, Marcus

CS Department of Neurology, Institute of Psychiatry, King's College London, London, SE1 1UL, UK

SO Journal of Neurochemistry (2001), 78(1), 13-23

CODEN: JONRA9; ISSN: 0022-3042 PB Blackwell Science Ltd.

DT Journal

LA English

OSC.G 13 THERE ARE 13 CAPLUS RECORDS THAT CITE THIS RECORD (13 CITINGS)
RE.CNT 50 THERE ARE 50 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 11 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN

TI In situ crosslinking of proteins for wound sealant

This fitt crosslinking of proteins for wound sealaim.

This invention relates to materials and methods for in situ crosslinking of proteins, including collagen, with peroxidase, including horseradish peroxidase, and H202 to form biocompatible semi-solid gels useful in a number of biol. and food product applications. The mixture applied to the wound sealing further comprises at least one addnl. agent selected from the group consisting of proteins, vaccine antigens, adjuvants, growth factors, microbeads and drugs, such as antimicrobials. The protein addnl. agent is selected from the group consisting of bovine serum albumin, fibrinogen, fibronectin, fibroblast growth factor, and human placental hyaluronic acid. A method of forming a semisolid crosslinked polymer on the surface of meat or poultry tissues for use as a food binding/restructuring agent comprises the steps of crosslinking a protein with a peroxidase in the presence of peroxide. Also, a method for growing dermal fibroblasts in vitro comprises the steps of growing the fibroblasts in a peroxide crosslinked collagen polymer.

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AN 2001:380339 HCAPLUS <<LOGINID::20091001>>
DN
    134:371845
TT
     In situ crosslinking of proteins for wound sealant
TN
    Miller, Douglas R.; Tizard, Ian R.; Keeton, Jimmy T.; Prochaska, Jerry F.
PA The Texas A & M University System, USA
SO PCT Int. Appl., 61 pp.
    CODEN: PIXXD2
    Patent
LA
    English
FAN.CNT 1
                     KIND DATE APPLICATION NO. DATE
ΡI
    WO 2001035882 A1 20010525 WO 2000-US31450 WO 2001035882 A9 20020815
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                         A1 20030102 EP 2000-979179
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US 1999-166024F P 19991115 <--
US 1909-16004 P 1999115 <--
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PRAI US 1999-165567P
ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OSC.G 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD (2 CITINGS)
RE.CNT 2
             THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 12 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
L5
TI
    Compounds for inhibiting diseases and preparing cells for transplantation
AB
    Methods and compns. are provided for inhibiting, preventing and treating
     amyloid depositions, e.g. in pancreatic islets, wherein the amyloidotic
     deposits are islet amyloid polypeptide (IAPP)-associated amyloid deposition
     or deposits. Accordingly, the compns. and method of the invention are
     useful for inhibiting IAPP-associated amyloidosis in disorders in which such
     amyloid deposition occurs, such an diabetes. The invention also provides
     a process for the preparation of cells suitable for transplantation into a
     mammal, which cells are capable of forming fibrils, said process
     comprising contacting the cells with an inhibitor of fibril formation.
     Also provided are a culture medium comprising the inhibitor and cells for
     transplantation. One example compound prepared was
     4-phenyl-1-(3-sulfopropyl)-1,2,3,6-tetrahydropyridine and its sodium salt.
     2001:50467 HCAPLUS <<LOGINID::20091001>>
AN
DN
    134:95503
     Compounds for inhibiting diseases and preparing cells for transplantation
IN
    Clark, Anne; Fraser, Paul; Verchere, Bruce; Gupta, Ajay; Migneault, David;
     Szarek, Walter; Weaver, Donald
    Isis Innovation Limited, UK; Neurochem, Inc.
PA
SO
    PCT Int. Appl., 62 pp.
    CODEN: PIXXD2
    Patent
LA English
FAN.CNT 1
     PATENT NO. KIND DATE APPLICATION NO. DATE
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PI WO 2001003680 A2 20010118 WO 2000-GB2623 20000707 <--
WO 2001003680 A3 20020711
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             HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
             LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
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                               20010118 CA 2000-2375628
20020911 EP 2000-946060
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                          A1
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         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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     US 20070015737 A1 20070118 US 2005-265537
                                                                   20051102 <--
PRAI GB 1999-16214
                         Α
                                19990709 <--
    GB 1999-16214 A 19990/09 <--
US 1999-16315 A 19990712 <--
WO 2000-GB2623 W 20000707 <--
US 2002-30350 B1 20021108 <--
ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OS MARPAT 134:95503
              THERE ARE 6 CAPLUS RECORDS THAT CITE THIS RECORD (6 CITINGS)
RE.CNT 9
              THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
L5
   ANSWER 13 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
TΙ
    Compositions and methods for treating amyloidosis
AB
    Therapeutic compds. and methods for modulating amyloid aggregation in a
     subject, whatever its clin. setting, are described. Amyloid aggregation
     is modulated by the administration to a subject of an effective amount of a
     therapeutic compound [(R1Zk)(R2Qm)N]pTYs [R1, R2 = H, (un)substituted alkyl,
     (un) substituted aryl; Z, Q = C(0), C(S), SO2, SO; k, m = 0, 1, with
     provisions; p, s = pos. integer such that biodistribution of therapeutic
     compound for intended target site is not prevented while maintaining
     activity of therapeutic compound; T = linking group; Y = AX; A = anionic
     group at physiol. pH; X = cationic group], or a pharmaceutically
     acceptable salt or ester, such that modulation of amyloid aggregation
     occurs. Preparation of e.g. 8-methoxy-5-quinolinesulfonic acid sodium salt is
     described.
AN
    2000:772432 HCAPLUS <<LOGINID::20091001>>
DN
    133:329624
ΤI
     Compositions and methods for treating amyloidosis
    Gordon, Heather; Szarek, Walter; Weaver, Donald; Kong, Xiangi
IN
PA
     Oueen's University at Kingston, Can.; Neurochem, Inc.
     PCT Int. Appl., 68 pp.
SO
     CODEN: PIXXD2
     Patent
DT
LA
     English
FAN.CNT 2
                        KIND DATE APPLICATION NO. DATE
     PATENT NO.
     WO 2000064420 A2 20001102
WO 2000064420 A3 20021107
                                          WO 2000-CA494
                                                                   20000428 <--
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             LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE,
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SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW

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     BR 2000010099
     EP 1276483
                            A2
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     EP 1276483
                            B1 20090902
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     JP 2003517458 T
                                  20030527 JP 2000-613411
                                                                            20000428 <--
     CN 1523992
                            A
                                   20040825
                                                CN 2000-809415
                                                                           20000428 <--
     CN 100482233 C 20070629 NZ 1919-5433 NZ 543319 A 20070629 NZ 1919-5433 NZ 543319 A 20094041 CN 2008-10149852 MX 2001010835 A 20030714 MX 2001-10835 NZ 825255 B1 20080416 KR 2001-713824 US 20040198832 A1 20041007 US 2003-639609 AU 2005202454 A1 20050630 AU 2005-202454 A1 20050630 AU 2005-202454 AI 20050630 NZ 2007-721102 NZ 2008-125842
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                                                                           20000524 <--
                                  20030714 MX 2001-10835
                                                                           20011025 <--
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                                   20041007 US 2003-639609
                                                                           20030811 <--
                                   20050630 AU 2005-202454
                                                                           20050603 <--
     AZ 200302434 A 20070927 KR 2007-721102 US 20080227967 A1 20080918 US 2008-125842 US 20080229759 A1 20081030 AU 2008-227580 AU 2008-229759 A1 20081030 AU 2008-229759 US 1999-131464P P 19990428 ---
                                                                           20070914 <--
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                                                                           20080707 <--
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PRAI US 1999-131464P
                          P
P
B1
     US 1999-135545P
                                    19990524 <--
     US 1999-133343F
US 1999-143123P
US 2000-560505
AU 2000-42824
                                   19990709 <--
                                    20000427
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                                    20000428 <--
     WO 2000-CA494
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     US 2000-576677 A
                                  20000523 <--
     AU 2000-49050
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                                  20000524 <--
     CN 2000-810720
KR 2001-713824
                           A3 20000524 <--
                           A3 20011029 <--
     US 2003-429198
                            A3 20030502
                            B1 20030811
     US 2003-639609
     AU 2005-203635
                            A3 20050815
ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OS MARPAT 133:329624
OSC.G 11 THERE ARE 11 CAPLUS RECORDS THAT CITE THIS RECORD (12 CITINGS)
RE.CNT 4
               THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD
               ALL CITATIONS AVAILABLE IN THE RE FORMAT
     ANSWER 14 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
TI Phosphocholine surfactants and their use
AB
   Disclosed are detergents or surfactants based on amphipathic
     phosphocholine compds. to improve pharmaceutical formulations and their
     use as pharmaceutical excipients.
     2000:553420 HCAPLUS <<LOGINID::20091001>>
AN
DN
     133:155464
TΙ
     Phosphocholine surfactants and their use
IN
     Morimoto, Bruce H.; Barker, Peter L.; Hernandez, Vincent; Piper, Cass K.
PA
     Amur Pharmaceuticals, Inc., USA
     PCT Int. Appl., 25 pp.
SO
     CODEN: PIXXD2
DT
     Patent
T. Z
     English
FAN.CNT 1
     PATENT NO. KIND DATE APPLICATION NO. DATE
     WO 2000045822 A1 20000810 WO 2000-US2395 20000128 <--
PΤ
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            TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG,
            KZ, MD, RU, TJ, TM
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     EP 1150685
                        A1 20011107 EP 2000-913304
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     JP 2002536335
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     US 6489369
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                                        US 2000-493359
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PRAI US 1999-118499P
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                              19990203 <--
     WO 2000-US2395
                       W
                              20000128 <--
ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OS MARPAT 133:155464
RE.CNT 5
             THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD
             ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 17 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
     Biocompatible composite material
    Biocompatible composites useful as a bone or tooth substitute material or
     for coating implants of metal, ceramic, Si, or plastics comprise an inorg.
     gel containing homogeneously embedded scleroproteins or their hydrolysis
     products and/or glycosaminoglycans. These composites promote the
     deposition of basic Ca phosphate phases and are hard, strong, and wear
     resistant. Thus, Si(OEt)4 10, 1,4-dioxane 40, and 0.01M HC1 20 mL were
    stirred for 20 h at room temperature to form a stable SiO2 soluble This sol 7
was
    mixed with H2O 7, 10% aqueous ZrO2 sol 2.3 mL, and 1% collagen type I solution
     g to provide a clear sol which was used for dip coating a Ti test piece.
     After drying, the coating had a Vickers hardness of 44. On immersion in
     simulated blood, the coated Ti induced deposition of basic Ca phosphate
    within 12 h.
    1999:624672 HCAPLUS <<LOGINID::20091001>>
    131:233590
    Biocompatible composite material
IN
   Brasack, Ingo; Boettcher, Horst; Kallies, Karl-Heinz
    Feinchemie G.m.b.H. Sebnitz, Germany; Kallies Feinchemie AG
    Ger. Offen., 6 pp.
    CODEN: GWXXBX
    Patent.
    German
FAN.CNT 1
     PATENT NO.
                      KIND
                              DATE APPLICATION NO. DATE
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    DE 19811900
                       A1 19990923 DE 1998-19811900
                                                               19980318 <--
    DE 19811900
PRAI DE 1998-19811900
OSC.G 5
                              20031211
                              19980318
OSC.G 5
             THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD (5 CITINGS)
RE.CNT 8
             THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD
             ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 18 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN
    Sphingolipid derivatives, their preparation, and their therapeutic use
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L5

L5

AB

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LA

TT Derivs. of sphingolipids (Markush included) are provided. The compds. are useful in the treatment of abnormal cell proliferation, including benign and malignant tumors, the promotion of cell differentiation, the induction of apoptosis, the inhibition of protein kinase C, and the treatment of

inflammatory conditions, psoriasis, inflammatory bowel disease as well as proliferation of smooth muscle cells in the course of development of plaques in vascular tissue. The invention also includes a method for triggering the release of cytochrome c from mitochondria that includes administering an effective amount of a sphingolipid or its derivative or

to a host in need thereof. Further, the invention provides a method for treating bacterial infections, including those that influence colon cancer and other disorders of the intestine, that includes administering an effective amount of one of the active compds. identified herein.

AN 1999:529160 HCAPLUS <<LOGINID::20091001>>

DN 131:165335

- TΙ Sphingolipid derivatives, their preparation, and their therapeutic use TN Liotta, Dennis C.; Merrill, Alfred H., Jr.; Keane, Thomas E.; Schmelz, Eva M.; Bhalla, Kapil N. PA Emory University, USA
 - PCT Int. Appl., 140 pp.
- SO CODEN: PIXXD2
- DТ Patent.
- LA English

FAN. CNT 1

	CIT	_																	
	PA:	TENT :	NO.			KIND DATE					APPL	ICAT	ION :	DATE					
PΙ	WO 9941266					A1 19990819													
		W:							BG,										
			ES,	FI,	GB,	GE,	HU,	IL,	IS,	JP,	KE,	KG,	KΡ,	KR,	ΚZ,	LK,	LR,	LS,	
			LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	
			SE,	SG,	SI,	SK,	TJ,	TM,	TR,	TT,	UA,	UG,	US,	UZ,	VN				
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	AU 9927644					A		1999	0830		AU 1	999-	2764		1	9990	212	<	
	AU	7658	09			B2		2003	1002										
	EP	1053	243			A1		2000	1122		EP 1	999-	9081		19990212 <				
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	US	6610	835			B1		2003	0826		US 1	999-	2492	11		1	9990	212	<
	AU	2003	2350	51		A1		2003	0911		AU 2	003-	2350	51		2	0030	814	<
	US	2004	0039	212		A1		2004	0226		US 2	003-	6478		20030825 <				
PRAI	US	1998	-745	36P		P		1998	0212	<-	-								
	AU	1999	-276	44		A3		1999	0212	<-	-								
	US	1999	-249	211		A1		1999	0212	<-	_								
	WO	1999	-US3	093		W		1999	0212	<-	_								

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OS MARPAT 131:165335

OSC.G 17 THERE ARE 17 CAPLUS RECORDS THAT CITE THIS RECORD (21 CITINGS) RE.CNT 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- ANSWER 23 OF 31 HCAPLUS COPYRIGHT 2009 ACS on STN L5
- TΙ Synthetic phosphopeptides for treating bone diseases
- AB Phosphopeptides which significantly reduce bone loss or weakening are provided. A method for treating or preventing any conditions associated with bone loss or weakening by administering the phosphopeptides by oral or injectable means is also provided. After age 35, bone mass, mineral content and mech. strength of the bone begin declining gradually. The relationship between bone mass and age is shown. Examples of prevention of bone loss in an osteoporosis model are given for peptides such as Pse-Gly-Pse-Gly-Pse-Gly (Pse = O-phosphoserine).

AN 1998:55543 HCAPLUS <<LOGINID::20091001>>

DN 128:110877

OREF 128:21617a,21620a

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Synthetic phosphopeptides for treating bone diseases
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IN Kumagai, Yoshinari; Otaka, Akira PA Big Bear Bio, Inc., USA

SO PCT Int. Appl., 45 pp. CODEN: PIXXD2

DT Patent

LA English FAN.CNT 2

211110112 2												* ^ > m									
						KIND DATE															
PI	WO	9800	156			A1 19980108				1	WO 1	997-	US11		19970630 <						
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			KR,	LK,	LT,	LV,	MD,	MK,	MN,	MX,	NO,	NZ,	PL,	SG,	SI,	SK,	TR,	UA,			
			UZ.	VN.	AZ.	BY.	KZ.	RU,	TJ.	TM											
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	US		674																		
			8661																		
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										AU 1997-35871						11	9970	630	<		
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			126					1999			PD 1	997_	0321	na		1.	9970	630	/		
			126					2004			31 I	,,,	JJ24	0,5			,,,,,,	050	`		
	LP									C.D.	O.D.	T. IT.			0.7	D.III					
			AT,																		
	JP	2001	.5034	52		Т		2001	0313		JP 1	998-	5043	99		1	9970	630	<		
	ΑT	2759	61			T		2004	1015	- 2	AT 1	997-	9324	09		1	9970	630	<		
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	JP	2004	10676	87		A				JP 2003-274414											
PRAI	US	1996	-675	031		A		1996	0703	<	-										
	JP	1998	-504	399		A3		1997	0630	<	-										
	WO	1997	-US1	1426		W		1997	0630	<	_										
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT